



187 00

NAVAL AIR STATION FORT WORTH JRB CARSWELL FIELD TEXAS

ADMINISTRATIVE RECORD COVER SHEET

File: $10^{12} + 10^{12} = 10^{12}$

12.

187 01

11-3517-0111

INSTALLATION RESTORATION PROGRAM (IRP) RCRA FACILITY INVESTIGATION

SAMPLING AND ANALYSIS PLAN

Carswell Air Force Base, Fort Worth, Texas

February 1994

Revised Final



PREPARED FOR

AIR FORCE BASE CONVERSION AGENCY (AFBCA/OL-H) CARSWELL AIR FORCE BASE, TEXAS 76127

UNITED STATES AIR FORCE AIR FORCE CENTER FOR ENVIRONMENTAL EXCELLENCE BASE CLOSURE RESTORATION DIVISION (HQ AFCEE/ESB) BROOKS AIR FORCE BASE, TEXAS 78235-5328 February 23, 1994

Air Force Center for Environmental Excellence HQ AFCEE/ESB 8001 Inner Circle Drive Suite 2 Brooks Air Force Base, TX 78235-5328

Attention:

Chris Hobbins (Team Chief)

Subject:

Carswell Air Force Base

Final Sample and Analysis Plan Contract No. F33615-90-D-4008

Delivery Order No. 0011 Law Project No. 11-3517-0111

Dear Mr. Hobbins:

Law Environmental, Inc., Government Services Division is pleased to submit the enclosed 25 copies of the Revised Final Sample and Analysis Plan to the Air Force Center for Environmental Excellence (AFCEE) for approval.

If you have questions or comments, please contact us at (404) 499-6800.

Sincerely,

John F. O'Brien

John 3. O Bren

Project Manager

E. Fred Sharpe, Jr., P.E.

E. Thed Shapp

Principal

REPORT DOCUMENTATION PAGE

Form Approved OMB No. 0704-0188

Public reporting burden for this collection of information is estimated average 1 hour per response, including the time for reviewing instructors, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect on this collection of information, including suggestions for reducing this burden to Washington Headquarters Services. Directorate for information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reduction Project (0704-0188), Washington, DC 20503.

1. AGENCY USE ONLY (Leave blank)	2. REPORT DATE	3. REPORT TYPE AND DATES COV	187 03
	FEBRUARY 1994	REVISED FINAL	18/ 33
4. TITLE AND SUBTITLE		5. FUNDING	GNUMBERS
INSTALLATION RESTORATION		CONT	RACT NO.
RCRA FACILITY INVESTIGAT			5-90-D-4008/0011
SAMPLING AND ANALYSIS F	PLAN		
6. AUTHOR(S)			
LAW ENVIRONMENTAL, INC	. (LAW)		
		Ì	
7. PERFORMANCE ORGANIZATION NAI	ME/S) AND ADDRESS/ES)	8 PERFOR	RMING ORGANIZATION
LAW ENVIRONMENTAL, INC	• • • • • • • • • • • • • • • • • • • •	_	NUMBER
GOVERNMENT SERVICES D		†	
114 TOWNPARK DR.			(N/A)
KENNESAW, GA 30144			
9. SPONSORING/MONITORING AGENCY	Y NAME(S) AND ADDRESS(ES)	10. SPONS	ORING/MONITORING
AIR FORCE CENTER FOR E	NVIRONMENTAL EXCELLENCE	AGENO	Y REPORT NUMBER
	ATION DIVISION (AFCEE/ESB)		(1)(4)
BROOKS AFB, TEXAS 78235		j	(N/A)
11. SUPPLEMENTARY NOTES		<u> </u>	
12a. DISTRIBUTION/AVAILABILITY STA	TEMENT	12b. DISTR	RIBUTION CODE
APPROVED FOR PUBLIC RE			
DISTRIBUTION UNLIMITED	,		
13. ABSTRACT (Maximum 200 words)			
, ,	., (LAW), WILL, INITIALLY, COND	HOT A DODA EACH ITV INIVE	STIGATION AT TWO SITES AT
	SE (CAFB), TEXAS, UNDER THE A		
	THE REQUIREMENTS OF IRP,		
	ESIGNED TO CHARACTERIZE E		
· · · ·	MEDIAL DESIGN AND REMEDIA		
	CLOSURE (BRAC) PROGRAM. TH		
	WMU 64) AND POL TANK FARM (
, : :::= : = ::	. ,	- · · · · · · · · · · · · · · · · · · ·	
			j
14. SUBJECT TERMS			15. NUMBER OF PAGES
RCRA FACILITY INVESTIGAT	TION		200
			16. PRICE CODE
17. SECURITY CLASSIFICATION	18. SECURITY CLASSIFICATION	19. SECURITY CLASSIFICATION	20. LIMITATION OF ABSTRACT
OF REPORT	OF THIS PAGE	OF ABSTRACT	UL
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	OL.

INSTALLATION RESTORATION PROGRAM (IRP) RCRA FACILITY INVESTIGATION FINAL SAMPLING AND ANALYSIS PLAN FOR

CARSWELL AFB FORT WORTH, TEXAS 76127-5000

FEBRUARY 1994

Prepared by:

Law Environmental, Inc. 114 TownPark Drive Kennesaw, Georgia 30144

CONTRACTOR CONTRACT NO. F33615-90-D-4008 DELIVERY ORDER NO. 0011

United States Air Force Air Force Center For Environmental Excellence Base Closure Restoration Division (HQ AFCEE/ESB) Brooks Air Force Base, Texas 78235-5328 Mr. Chris Hobbins (Team Chief)

This Sampling and Analysis Plan (SAP) has been developed for the RCRA Facility Investigation (RFI) activities at Carswell Air Force Base (Carswell AFB). Procedures outlined in this plan are designed to describe the collection of geologic data, hydrologic data, environmental samples, laboratory analysis of those samples for potential contaminants, evaluation of the analytical results and field measurements with respect to quality control data and the interpretation and analysis of QA/QC reviewed data. The plan will be effective after final approval.

The success of Carswell AFB's Installation Restoration Program depends on team effort and total dedication from parties involved. Therefore, efforts will be focused on achieving and maintaining compliance with this Sampling and Analysis Plan and pertinent regulations.

The point of contact for this investigation is as follows:

Mr. Chris Hobbins
Team Chief (TC)
HQ AFCEE/ESB
8001 Inner Circle Drive
Suite 2
Brooks AFB, Texas 78235-5328
Phone: (210) 536-5261

This Sampling and Analysis Plan has been prepared for the United States Air Force by Law Environmental, Inc. for the purpose of aiding in the implementation of a final remedial action plan under the Air Force Installation Restoration Program (IRP). report relates to actual or possible releases of potentially hazardous substances, its release prior to an Air Force final decision on remedial action may be in the public's interest. limited objectives of this plan and the ongoing nature of the IRP, along with the evolving knowledge of site conditions and chemical effects on the environment and health, must be considered when evaluating this report, since subsequent facts may become known which may make this plan premature or inaccurate. Acceptance of this sampling and analysis work plan in performance of the contract under which it is prepared does not mean that the Air Force adopts the conclusions, recommendations or other views expressed herein, which are those of the contractor only and do not necessarily reflect the official position of the United States Air Force.

Copies of this plan may be purchased from:

Government agencies and their contractors registered with the Defense Technical Information Center (DTIC) or should direct their requests for copies of this work plan to:

Defense Technical Information Center Cameron Station Alexandria, VA 22304-6145

Non-government agencies may purchase copies of this document from:

National Technical Information Service (NTIS) 5285 Port Royal Road Springfield, VA 22161

3517-0111.10

SAMPLING AND ANALYSIS PLAN (SAP) PREFACE

Law Environmental, Inc. (Law) was contracted by the U.S. Air Force Center for Environmental Excellence (AFCEE) to perform a RCRA Facility Investigation (RFI) at two sites at Carswell AFB, Texas. The two sites to be investigated include: Unnamed Stream (IRP Site SD-13/SWMU 64) and POL Tank Farm (IRP Site ST-14/SWMU 68). The primary objective of this field investigation is to investigate the extent of soil and ground water contamination at each site and assess the overall environmental status of the sites in order to support the recommendation of appropriate further actions. Project objectives will be achieved through the use of the following methods of investigation: geophysical and geochemical surveys; soil and ground water samples for field screening and laboratory analysis; and surface water and sediment samples for laboratory analysis.

The SAP outlines the site objectives, the data quality objectives, the field activities, sample collection, and laboratory analytical procedures required for the RFI at Carswell AFB.

The SAP is composed of two documents - the Quality Assurance Project Plan (QAPP) and the Field Sampling Plan (FSP).

The QAPP consists of detailed information on defining and assuring that the Data Quality Objectives (DQOs) are achieved. DQOs are considered through various project tasks, including writing of plans, field work, and laboratory analysis. The QAPP delineates the procedures necessary to achieve DQO goals.

The FSP describes field tasks necessary for implementing the project objectives. Field tasks are described in detail to ensure that the DQOs are achieved during field activities.

Mr. John O'Brien is the Project Manager for the RFI. Members of the field investigation team will be selected prior to commencement of field activities.

The QAPP was prepared by Ms. Darice Kurtzer and reviewed by Mr. Jerry Preston and Mr. Fred Sharpe. The FSP was prepared by Mr. Jim Beaver and Ms. Darice Kurtzer and reviewed by Mr. Jerry Preston and Mr. Fred Sharpe.

The support and assistance of Mr. Chris Hobbins (AFCEE) and personnel at Carswell AFB are greatly appreciated.

John F. O'Brien

Project Manager

E. Fred Sharpe, Mr., P.E

Principal

Louis S. Karably, P.E.

Program Manager

TABLE OF CONTENTS

		<u>Page</u>
LIST	OF AC	PRONYMS
1.0	QUAL	TY ASSURANCE PROJECT PLAN
	1.1	INTRODUCTION
		1.1.1 The U.S. Air Force Installation Restoration
		Program
	1.2	PROJECT DESCRIPTION
		1.2.1 Project Background 1-4
		1.2.2 Project Scope and Objectives 1-6 1.2.3 Subcontractors
	1.3	PROJECT ORGANIZATION AND RESPONSIBILITIES 1-14
		1.3.1 Project Organization
		1.3.3 Project QA Responsibilities 1-21
	1.4	QUALITY ASSURANCE OBJECTIVES FOR MEASUREMENT DATA 1-22
		1.4.1 Definitions of QA Criteria 1-23
		1.4.2 Goals 1-26
	1.5	SAMPLING PROCEDURES
		1.5.1 Sampling Protocols 1-29
		1.5.2 Sample Handling 1-29
	1.6	SAMPLE CUSTODY
		1.6.1 Field Operations 1-30
		1.6.2 Laboratory Operations 1-30
	1.7	FIELD EQUIPMENT CALIBRATION PROCEDURES 1-30
	1.8	ANALYTICAL PROCEDURES
		1.8.1 Identification of Methods 1-33
		1.8.2 Detection Limits and Quantitation Criteria 1-3
		1.8.3 Method Calibration 1-3

TABLE OF CONTENTS (Continued)

		<u>'</u>	<u>Page</u>
	1.9	DATA REDUCTION, VALIDATION, AND REPORTING	1-60
		1.9.1 Data Management	1-60 1-60 1-60 1-66
	1.10	INTERNAL QUALITY CONTROL CHECKS	1-66
		1.10.1 Field Quality Control	1-66 1-66 1-77
	1.11	PERFORMANCE AND SYSTEM AUDITS	1-78
		1.11.1 Systems Audits	1-78 1-79 1-80
	1.12	PREVENTIVE MAINTENANCE	1-80
	1.13		1-80
		1.13.1 Formulas	1-87
	1.14	CORRECTIVE ACTION	1-88
		1.14.1 Response	1-89
	1.15	QUALITY ASSURANCE REPORTS	1-90
		1.15.1 Reporting Procedure	1-91 1-91
2.0	FIEL	D SAMPLING PLAN (FSP)	2-1
	2.1	FIELD OPERATIONS	2-1
		2.1.1 Site Reconnaissance, Preparation and Restoration	2-1 2-5

187 11

TABLE OF CONTENTS (Continued)

		<u>Page</u>
	2.1.3 Geochemical Survey	. 2-7
	2.1.4 Borehole Construction, Lithologic Sampling	
	and Logging	. 2-8
	2.1.5 Monitoring Well Construction and	
	Installation	2-10
	2.1.6 Aquifer Tests (Slug Tests)	2-17
	2.1.7 Surveying	2-20
	2.1.8 Equipment Decontamination	2-21
	2.1.9 Waste Handling	2-22
	2.1.10 Field Activity Summary	2-23
2.2	ENVIRONMENTAL SAMPLING	2-23
	2.2.1 Procedures for Collection of Samples	2-23
	2.2.2 Sample Handling	2-23
	2.2.2 Sample Handling	2-41
	2.2.4 QC Samples	2-40
	2.2.5 Sample Analysis Summary	2-50
	2.2.5 bample Analysis bundlary	2-50
2.3	FIELD MEASUREMENTS	2-50
	2.3.1 Parameters for Field Characterization of	
	Samples	2-50
	2.3.2 Equipment Calibration	2-51
	2.3.3 Equipment Maintenance	2 - 52
	2.3.4 Decontamination of Field Instruments	2-52
2.4	FIELD QA/QC PROGRAM	2-53
	2.4.1 Control Parameters	2-53
	2.4.2 Control Limits and Corrective Actions	2-53
2.5	RECORD KEEPING	2-53
2.6	SITE MANAGEMENT	2-54
REFERENCE		. R-1

LIST OF FIGURES

<u>Figur</u>	<u>re</u>	<u>Page</u>
1-1	Location Map of Carswell AFB	. 1-3
1-2	Location of East Area Sites	. 1-5
1-3	Locations of French Drain and Oil/Water Separator, Site SD13	. 1-7
1-4	Proposed Geochemical Sampling Area	1-12
1-5	Project Organization and QM Responsibilities	1-16
2-1	Location of East Area Sites	. 2-2
2-2	Location of Wells and Surface Water Sampling Locations and Approximate Locations of French Drains and Oil/Water Separator	. 2-3
2-3	Proposed Geochemical Sampling Area	. 2-4
2-4	Cone Penetrometer Sampler Diagram	. 2-9
2-5	Soil Test Boring Record	2-11
2-6	Monitoring Well Installation Diagram	2-13
2-7	Well Development Data	2-18
2 - 8	Request for Analytical Form	2-46
2-9	Chain of Custody Record	2-48
2-10	Daily Quality Control Report	2-55

LIST OF TABLES

<u>Table</u>	2	<u>Page</u>
1-1	Potential Contaminants RCRA Facility Investigation	. 1-9
1-2	Parameters of Interest by Site	1-10
1-3	Analytical Levels	1-27
1-4	Analytical Requirements - Soil and Sediment Samples .	1-32
1-5	Analytical Requirements - Ground and Surface Water Samples	1-34
1-6	Project-Required Reporting Limits - Soil/Sediment	1-38
1-7	Project-Required Reporting Limits - Ground Water/Surface Water	1-41
1-8	DFTPP Key Ions and Abundance Criteria	1-50
1-9	BFB Key Ions and Abundance Criteria	1-51
1-10	Volatile Internal Standards with Corresponding Analytes Assigned for Quantitation	1-53
1-11	Semi-Volatile Internal Standards with Corresponding Analytes Assigned for Quantitation	1-55
1-12	Data Evaluation Categories	1-63
1-13	Data Qualification Flags	1-65
1-14	Analysis Summary: Sub-Surface Soil From Borings	1-68
1-15	Analysis Summary: Ground Water - Monitoring Wells	1-69
1-16	Analysis Summary: Ground Water - Field Screen	1-71
1-17	Analysis Summary: Surface Water	1-72
1-18	Analysis Summary: Sediment	1-73
1-19	Routine Maintenance Schedule for GC	1-81
1-20	Routine Maintenance Schedule for GC/MS	1-82
1-21	Routine Maintenance Schedule for ICP	1-83

187 14

LIST OF TABLES (Continued)

<u>Tabl</u>	<u>e</u>	<u>Page</u>
1-22	Routine Maintenance Schedule for AA Spectrophotometer	1-84
1-23	Routine Maintenance Schedule for IR Spectrophotometer	1-85
2-1	Field Tasks - RCRA Facility Investigation	2-24
2-2	Field Equipment Checklist	2-25
2-3	Sampling Summary: Sub-Surface Soil from Borings	2-28
2-4	Sampling Summary: Ground Water - Monitoring Wells	2-29
2-5	Sampling Summary: Ground Water - Field Screening	2-31
2-6	Sampling Summary: Surface Water	2-32
2 - 7	Sampling Summary: Sediment	2-33
2-8	Sample Containers, Amounts, and Preservation - Soil and Sediment Samples	2-42
2-9	Sample Containers, Amounts, and Preservation - Ground-Water and Surface Water Samples	2-43

LIST OF APPENDICES 187 15

APPENDIX A	QUALITY CONTROL REQUIREMENTS AND REPORTING LIMITS
APPENDIX B	QUALITY ASSURANCE PROGRAMS FOR CHEMICAL ANALYSIS LABORATORIES
APPENDIX C	PHOTOIONIZATION DETECTOR
APPENDIX D	ph meter
APPENDIX E	SPECIFIC CONDUCTIVITY AND TEMPERATURE METER
APPENDIX F	TURBIDIMETER
APPENDIX G	MILLIPORE OPERATION AND MAINTENANCE INSTRUCTIONS

LIST OF ACRONYMS

Ambient Condition Blank AB ABB-ENV ABB Environmental Services, Inc. AFB Air Force Base AFCEE Air Force Center for Environmental Excellence ARAR Applicable or Relevant and Appropriate Requirements ASTM American Society for Testing and Materials Corrective Action CA CF Calibration Factor Continuing Calibration Blank CCB CCC Continuing Calibration Check CCV Continuing Calibration Verification CE Civil Engineering CERCLA Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (PL-96-510) - SUPERFUND CES Civil Engineering Squadron CEO Chief Executive Officer CFR Code of Federal Regulations CLP Contract Laboratory Program CV Coefficient of Variation CVAA Cold Vapor Atomic Absorption DC Drill Cuttings Department of Transportation DOT DOCR Daily Quality Control Report DOO Data Quality Objectives Equipment Blank (Rinsate) EB EMSL Environmental Monitoring Systems Laboratory FS Feasibility Study FSP Field Sampling Plan GC/MS Gas Chromatography/Mass Spectrometry GFAA Graphite Furnace Atomic Absorption

Photoionization Detector (trade name)

Initial Calibration Blank

Sample Identification

HNu ICB

ID

LIST OF ACRONYMS (Continued)

ICP Inductively Coupled Plasma

IRP Installation Restoration Program

IRPIMS Installation Restoration Program Information Management

System

ITIR Informal Technical Information Report

Law Environmental, Inc., Government Services Division

LEGS Law Environmental, Inc. - Government Services Division

LENL Law Environmental National Laboratories

LENL-P Law Environmental National Laboratories - Pensacola,

Florida

LCS Laboratory Control Standards

MCL Maximum Contaminant Level

MDL Method Detection Limit

MITRE MITRE Corporation

MQL Maximum Quantitation Limit

MS Matrix Spike

MSD Matrix Spike Duplicate

MW Monitoring Well

NCP National Contingency Plan

NTU Nephelometric Turbidity Unit

OSWER Office of Solid Waste and Emergency Response

OVA Organic Vapor Analyzer

PA Preliminary Assessment

PARCC Precision Accuracy Representativeness Completeness

Comparability

PE Professional Engineer

PID Photoionization Detector

POC Point of Contact

QA Quality Assurance

QA/QC Quality Assurance/Quality Control

QAMS Quality Assurance Management Staff

QCCS Quality Control Check Samples

LIST OF ACRONYMS (Continued)

QAPP Quality Assurance Project Plan

QC Quality Control

RAGS Risk Assessment Guidance for Superfund RCRA Resource Conservation and Recovery Act

RF Response Factor

RI Remedial Investigation

RLS Registered Land Surveyors

RPD Relative Percent Difference

RRF Relative Response Factor
RRT Relative Retention Time

RSD Relative Standard Deviation

RT Retention Time

SAP Sampling and Analysis Plan

SB Soil Boring

SC Specific Conductance

SD Surface Sediment
SI Site Investigation

SOP Standard Operating Procedures

SPCC System Performance Check Compound

SPT Standard Penetration Test

SW Surface Water

TB Trip Blank
TC Team Chief

TEG TransGlobal Environmental Geochemistry, Texas

TPM Technical Project Manager
TWC Texas Water Commission
USAF United States Air Force

USEPA United States Environmental Protection Agency

USGS United States Geological Survey

1.1 INTRODUCTION

This RCRA Facility Investigation (RFI) Quality Assurance Project Plan (OAPP) is provided in support of the United States Air Force (USAF) Installation Restoration Program (IRP) for two sites located Carswell Air Force Base (Carswell AFB), investigation will adhere to those procedures specified by the RFI Guidance EPA 530/SW-89-031 and Provision VIII RCRA Facility Investigation of the permit issued to Carswell AFB by the Texas Water Commission (TWC) on February 7, 1991. Only the parameters of concern (Section 1.2.2) will be investigated; there is no evidence that other chemicals listed in 40 CFR 264 Appendix IX have ever been used at or adjacent to the two sites. This investigation will be supplemented by the Air Force Center for Environmental Excellence (AFCEE) in the Handbook to Support the Installation Restoration Program (IRP) Statements of Work: Volume I - Remedial Investigation/Feasibility Studies (RI/FS) (AFCEE, 1991).

1.1.1 The U.S. Air Force Installation Restoration Program

The objective of the Air Force IRP is to assess past hazardous waste disposal and spill sites at Air Force installations and to develop remedial actions consistent with the National Contingency Plan (NCP) for those sites which present a threat to human health and welfare or the environment. Additional information regarding the Air Force IRP is discussed in Section 1.0 of the Work Plan.

1.1.2 Purpose and Scope

The Sampling and Analysis Plan (SAP) describes those procedures necessary to perform a quality controlled sampling effort during

the execution of the project's scope of work. The activities addressed by the SAP include:

- · Installation of monitoring wells
- · Soil borings
- Acquisition of ground-water, surface and subsurface soil, surface water and sediment samples
- · Decontamination of equipment
- · Calibration of field instruments
- · Aquifer (slug) testing
- Maintenance of laboratory and field QA/QC
- · Geophysical survey
- · Chemical analysis
- · Preventative maintenance
- · Data quality assessment
- · Corrective action
- · Quality control (QC) checks
- · Quality assurance (QA) audits and reports

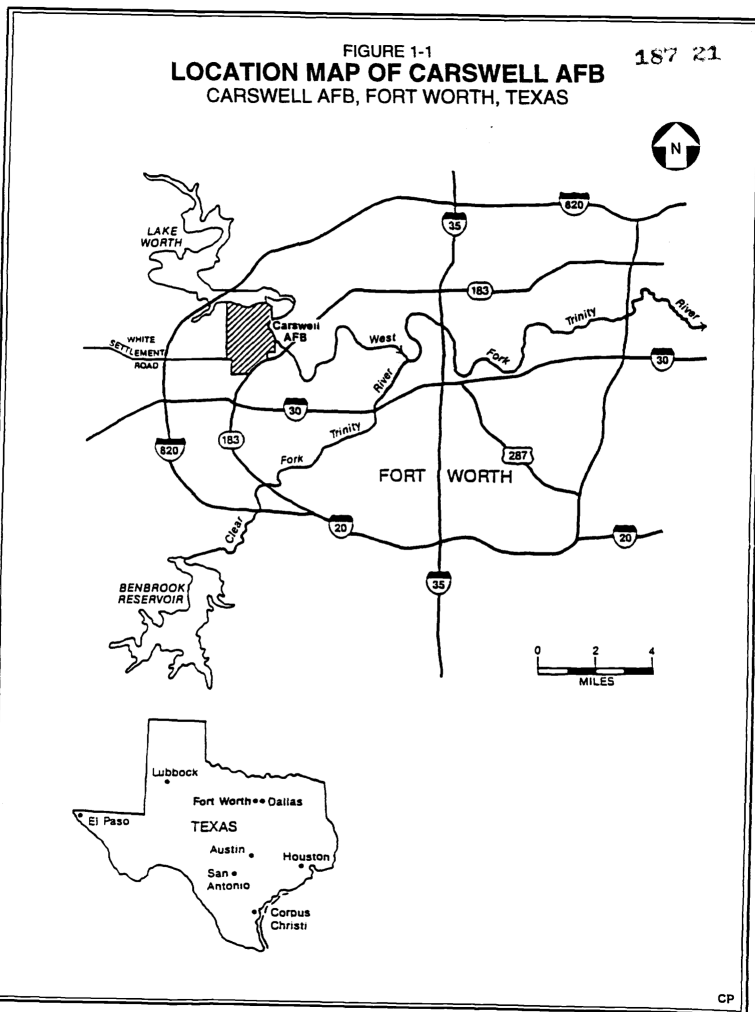
Specific quality control requirements are specified in Appendix A.

1.2 PROJECT DESCRIPTION

Carswell AFB is located in northeastern Texas, in Tarrant County, six miles west of downtown Fort Worth, as shown in Figure 1-1. This RFI for Carswell AFB will investigate the two sites listed below:

- Unnamed Stream (IRP Site SD-13/SWMU 64)
- Petroleum, Oil, Lubricants (POL) Tank Farm (IRP Site ST-14/SWMU 68)

Detailed site maps are presented in Section 2.1. The RFI for each site is designed to evaluate the presence and extent of



contamination. The following sections present the project background, scope and objectives.

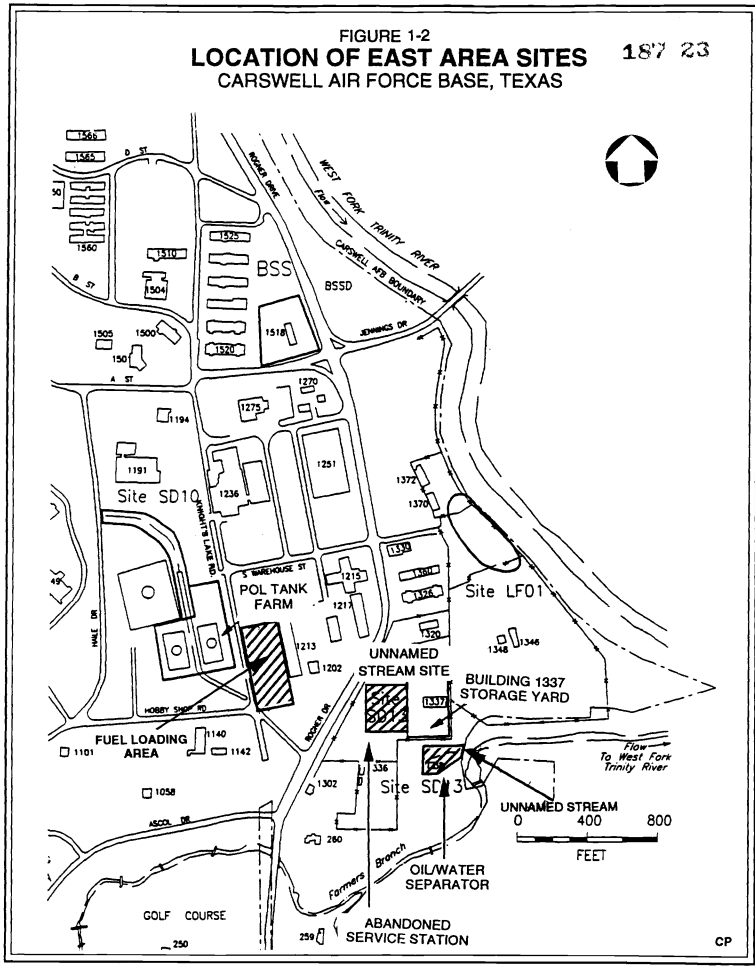
1.2.1 Project Background

The two sites included under this RFI were identified as possibly contaminated sites through the efforts of the Installation Restoration Program Phase I Records Search Report (CH2M Hill, 1984). Information from that report is supplemented by information from the literature and from the general findings of studies conducted by the Radian Corporation (Radian, 1986, 1991).

1.2.1.1 <u>POL Tank Farm Site (IRP Site ST-14/SWMU 68)</u> - The POL Tank Farm is located in the east area of Carswell AFB, west of and adjacent to Knight's Lake Road and north of Hobby Shop Road (Figure 1-2). The Fuel Loading Area is east of Knight's Lake Road.

Three aboveground POL storage tanks currently are in place at this site and an additional three tanks have been removed from the site. Leaking underground POL lines are suspected to have released an undetermined amount of fuel products into the soil and ground water at and south of the POL Tank Farm during the early 1960s. The leaking lines were reportedly located and replaced and no further fuel releases were documented after 1965. Previous studies (Radian, 1986, 1991) have found evidence of contamination by organic compounds, including ethylbenzene, benzene, chlorobenzene and total xylenes. Lead and chromium in excess of Maximum Containment Levels (MCLs) were also detected in the ground water at this site.

1.2.1.2 <u>Unnamed Stream Site (IRP Site SD-13/SWMU 64)</u> - The Unnamed Stream and Abandoned Service Station are located near the eastern boundary of the base, east of Rogner Drive and north of Farmer's



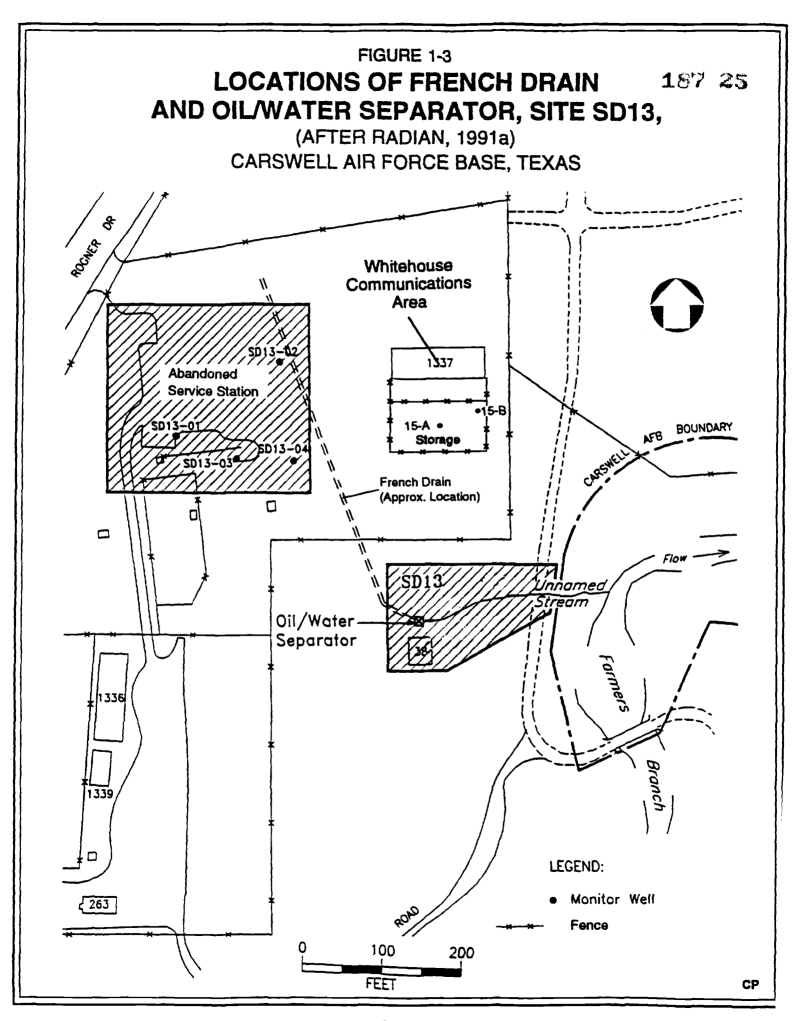
Branch. This site consists of two locations, the paved area around the Abandoned Service Station and the intermittent stream flowing from an oil/water separator to Farmer's Branch (Figure 1-3). The Abandoned Service Station area may contain one or more USTs that may have leaked petroleum products. The Unnamed Stream flows from the oil/water separator that is the terminus of a french drain

that may have leaked petroleum products. The Unnamed Stream flows from the oil/water separator that is the terminus of a french drain system installed in 1965, reportedly to recover fuel leaking from either the Abandoned Service Station USTs and/or the POL Tank Farm. Previous studies (Radian, 1986, 1991) detected low levels of organic contamination in the ground water at this site. Surface water samples contaminated with lead and arsenic in excess of MCLs were collected from the Unnamed Stream downstream from the oil/water separator.

1.2.2 Project Scope and Objectives

The following sections describe the site-specific scope and objectives. The objectives described below will be accomplished through the collection of geological data, hydrogeological data, and collection of environmental samples; the laboratory analysis of these samples for potential contaminants; the evaluation of the analytical results and field measurements with respect to quality control data; and the interpretation and analysis of valid data.

The data generated by this project must be of sufficient quality and quantity to meet the overall project objective, which is to improve site characterizations that would ultimately lead to remediation of the sites. The purpose of this RFI at Carswell AFB is to collect assessment data for ground water at POL Tank Farm and for soils, surface water, sediment, and ground water at the Unnamed Stream. This RFI will be prepared to meet requirements included in Part VIII of the RCRA Permit Number HW-50289, Attachment B. The focus of this work will be to characterize the spatial distribution of contamination at each site and its potential for transport. The



findings of this RFI will be used to support the recommendation of appropriate further actions. The criteria established in the Sampling Analysis Plan Will be used to determine the possible need for a corrective measures study to define and evaluate remedial action alternatives.

Health and Safety data will be used to establish the level of protection needed for the work party and other site-related personnel. These data will be gathered by the use of geophysical instruments, organic vapor analyzers (HNu), Draeger tubes, tri-gas monitors, and explosimeters utilized during non-intrusive and intrusive activities.

Table 1-1 presents a combined list of the potential contaminants at both sites. This table is compiled from the list of compounds in the SW846 EPA Test Methods for Evaluating Solid Waste. compounds listed are based upon the parameters of interest for each of the sites at Carswell AFB. In order to further evaluate the sites, samples of surface and subsurface soil, ground water, surface water, and sediment will be collected. The parameters of interest for each site for each of these matrices are presented in Table 1-2. Other parameters listed in 40 CFR 264 Appendix IX are not included as parameters of interest for these sites since there is no evidence of these chemicals ever being used at or adjacent to these sites. The number of samples to be collected are presented in Section 2.2. The collection and analysis of background soil and ground-water samples will be performed according to the AFCEE request on October 28, 1993. This AFCEE directive requested that background ground-water and soil samples be collected from the proposed upgradient monitoring well adjacent to Rogner Drive. following subsections describe the background scope and objectives for each site.

+ = Aromatic

3517-0111.10

TABLE 1-1

POTENTIAL CONTAMINANTS RCRA Facility Investigation Carswell Air Force Base, Texas

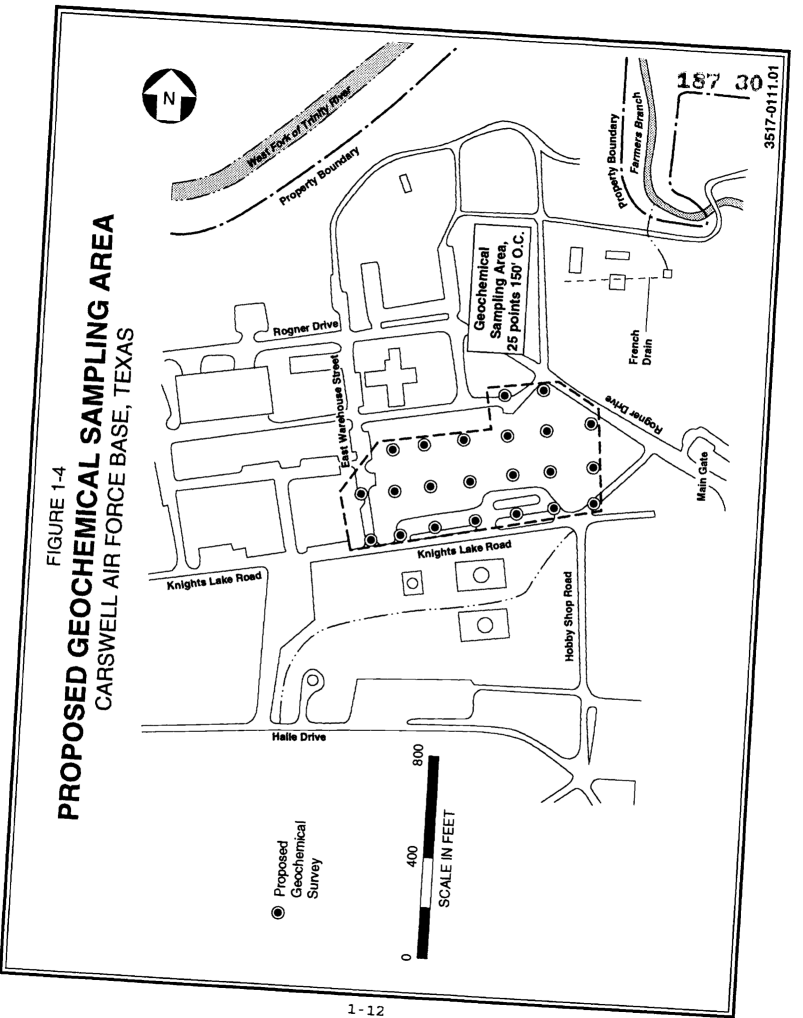
<u>Total Recoverable Petroleum Hydrocarbons</u> Oil Grease	Purgeable Aromatic Volatiles Benzene Chlorobenzene 1,2-Dichlorobenzene 1,4-Dichlorobenzene Ethylbenzene Toluene Xylenes Nonhalogenated Volatile Organics Diethyl ether Ethanol Methyl sthyl ketone (MEK) Methyl isobutyl ketone (MIBK) Oil Grease Grease	187
Metals (24) Aluminum Antimony Arsenic	Arsenic Barium Cadmium Calcium Chromium Cobalt Copper Iron Lead Magnesium Magnese Molybdenum Mercury Nickel Potassium Selenium Selenium Salver Sodium Thallium Vanadium	
Volatile Organics (34) Acetone Benzene + Bromodichloromethane	Bromonethane Bromomethane Bromomethane 2-Butanone (MEK) Carbon Disulfide Carbon tetrachloride Chlorobenzene Chloroethane Chloromethane 1,1-Dichloroethane 1,2-Dichloroethene 1,2-Dichloropropene trans-1,2-Dichloropropene trans-1,3-Dichloropropene Ethylbenzene + 2-Hexanone Methylene chloride 4-Methyl-2-pentanone Styrene 1,1,2,2-Tetrachloroethane Tetrachloroethene	1,1,2-Trichloroethane Trichloroethene Vinyl acetate Vinyl chloride Xylenes +

TABLE 1-2

PARAMETERS OF INTEREST BY SITE RCRA Facility Investigation Carswell Air Force Base, Texas

			ORGANIC				=	INORGANIC			OTHER
	AROMATIC VOLATILE		TOTAL RECOVERABLE		ol ol	SCREEN					
NO SITE NAME		VOCATILE VOCATILE		PEIROLEUM	ا ا کا ا	FOH 24					
	COMPANION	CHGANICS	HIDHOCAHBONS	HYDHOCAMB. GHEASE	GHEASE	METALS	ARSENIC	ARSENIC MERCURY LEAD SELENIUM	LEAD	SELENIUM	TCL
GROUND WATER / HYDROPUNCH SURVEY	ROPUNCH SU	IRVEY									
1 Unnamed Stream		×	×		*	>	>	>	>	>	>
2 POL Tank Farm	×		:	×	<	<	<	<	< ×	<	<
SURFACE WATER											
1 Unnamed Stream	×	×	×		×						
SUB~SURFACE SOIL											
1 Unnamed Stream	×	×	×								×
SEDIMENT											
1 Unnamed Stream						×					

- 1.2.2.1 <u>POL Tank Farm Site</u> Field tasks to be performed at the POL Tank Farm site include a geophysical survey and geochemical survey.
- A. <u>Geophysical Survey</u> A surface geophysical survey will be performed to help characterize the near surface conditions and to locate utilities and abandoned tanks and pipechases. A single type of geophysical survey (ground penetrating radar) will be performed at the POL Tank Farm site during the field work.
- B. Geochemical Survey A geochemical survey will be conducted at the POL Tank Farm site as part of the RCRA facility investigation effort. An estimated 25 survey points will be performed utilizing the Strataprobe™ System, which is a direct push method for ground-water investigation developed by Transglobal Environmental Geochemistry (TEG). These ground-water samples will provide supplemental analytical information in order to delineate the extent of contamination (see Figure 1-4).
- 1.2.2.2 <u>Unnamed Stream</u> Field tasks to be performed at the Unnamed Stream include a geophysical survey, monitoring well installation, aquifer testing, soil sampling and surface water/sediment sampling.
- A. <u>Geophysical Survey</u> A surface geophysical survey will be performed at both sites to help characterize the near surface conditions and to locate utilities and abandoned tanks and pipechases. A single type of geophysical survey (ground penetrating radar or other means) will be performed at the Unnamed Stream during the field work.



B. <u>Soil Sampling</u> - Two soil samples will be collected from each of three soil borings. The samples will be screened by hand held portable organic vapor analyzer(s) (OVAs) used to screen the air vapors at the head of the augers, in the breathing zone and in the soil sample headspace. The OVAs will be equipped with photoionization detectors (PID). The PID equipment manual is presented in Appendix C. One soil sample will be collected at the depth that has the highest OVA reading. The second sample will be collected at the groundwater interface. If the OVA screen results are nondetect, a default depth of five feet will be sampled.

In addition to the two sample collection points described above, two more soil samples will be collected from the soil boring located northwest of the French Drain and east of Rogner Drive; a total of four soil samples will be taken from this boring (see Figure 2-2). One surface soil sample will be collected at the 6- to 12-inch interval. The second and third samples will be collected at the depths with the two highest OVA readings, and the fourth sample will be collected at the ground-water interface. This upgradient soil boring will be completed as an upgradient monitoring well and this location will be used to establish background soil and ground-water data for the site.

Soil borings will be performed to allow installation of each monitoring well. The soil borings will provide subsurface information for well design and site stratigraphy as well as assessment of the presence of site constituents in surface and subsurface soils.

C. <u>Monitoring Well Installation</u> - Each of the three soil borings discussed above will be completed as shallow monitoring wells to assess the presence of site constituents in subsurface soils and the surficial aguifer. One monitoring well will be

located upgradient of this site and will be used to establish background information for this site. Two monitoring wells will be located downgradient of this site.

- D. Aquifer Testing In-situ hydraulic conductivity tests will be performed on the new monitoring wells a minimum of 24 hours after ground-water sampling. The tests will be conducted to determine formation permeability. The results of the tests will be used to estimate ground-water flow rates.
- E. <u>Surface Water/Sediment Sampling</u> Three surface water/sediment samples will be collected at this site to assess if site constituents are affecting surface waters and underlying sediments in the area.

1.2.3 Subcontractors

The subcontractors to be utilized for this RFI investigation along with their service to be performed are identified in Section 1.3.

1.3 PROJECT ORGANIZATION AND RESPONSIBILITIES

Project organization and responsibilities for the RFI for the two sites at Carswell AFB are discussed in the following sections.

1.3.1 Project Organization

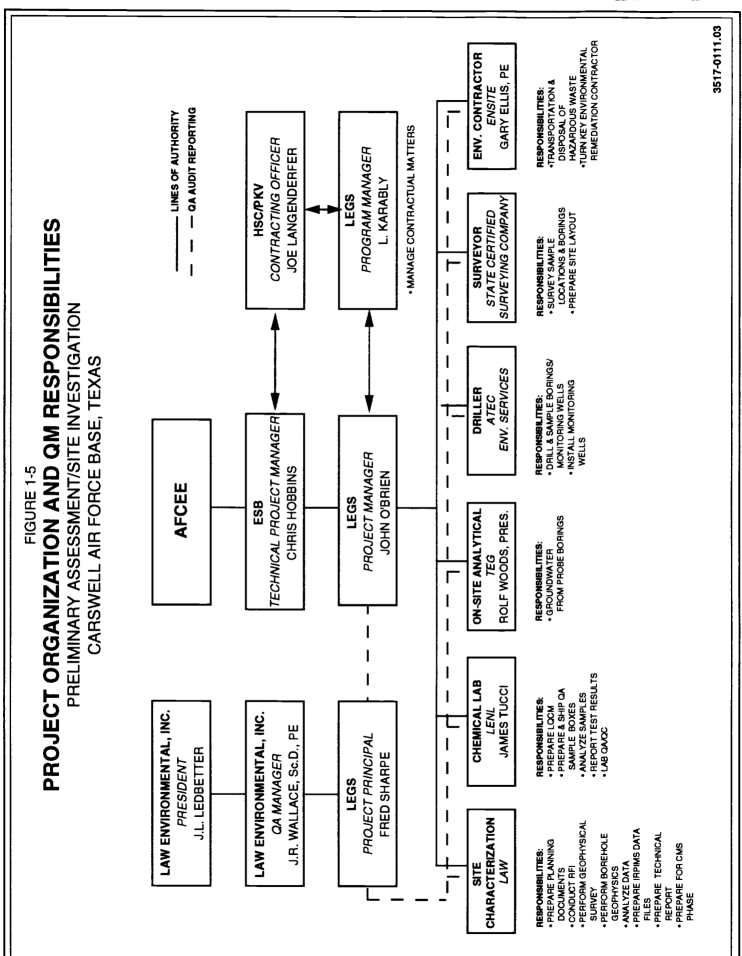
Law Environmental, Inc., Government Services Division (Law) will manage the project and provide services related to field samples, geophysics, data analysis, site characterization, and reporting.

ATEC Environmental Services of Dallas, Texas, will be subcontracted to perform the drilling services.

- Law Environmental National Laboratories Pensacola, Florida, (LENL-P) will be subcontracted as the primary laboratory to perform the chemical analysis of the soil and water samples.
- Transglobal Environmental Geochemistry (TEG) of Austin, Texas, will be subcontracted to perform the ground-water reconnaissance technique at the POL Tank Farm area. TEG will perform a ground-water field screening utilizing a direct push cone penetrometer type sampling system. The TEG sampling technique, Strataprobe™ is described in Section 2.1.3. They will also perform on-site analysis of the ground water collected. The TEG QA information is in Appendix B-2.
- An OSHA-certified and Texas state licensed local surveying company will be subcontracted to survey the soil borings, monitoring wells, and other sampling locations. In addition, the surveyor will prepare a site map.
- Ensite, Inc., will be subcontracted to handle the transport and disposal of drummed soil cuttings, decontamination fluids, and development and purge water generated during investigative activities.

The project organization is shown on Figure 1-5.

1.3.1.1 <u>Law Environmental, Inc.</u> - Law Environmental's Government Services Division will provide the project management, engineering and analysis, and sampling through its in-house resources. It is a branch of Law Environmental, Inc., which, in turn, is a whollyowned subsidiary of Law Companies Group, Inc. Law Companies Group, Inc., is entirely employee owned.



- 1.3.1.2 <u>Drilling Subcontractor</u> ATEC Environmental Services, Dallas, Texas will be responsible for drilling soil borings and installation of monitoring wells. The drilling engineer will follow direction from the Site Manager during field activities and from the Project Manager prior to and after field work.
- 1.3.1.3 Chemical Analysis Laboratories The Law Environmental, Inc., National Laboratories facility in Pensacola, Florida (LENL-P), is the Law Environmental, Inc., chemical testing laboratory which was established in 1989. LENL-P will be the laboratory responsible for providing sample shipping containers, chain-ofcustody documents, chemical analysis, reporting, and laboratory LENL-P will perform all analyses of soil, ground water, OA/OC. surface water and sediments, and associated QC samples. will report directly to the Project Manager during the project. LENL-P has integrated Quality Assurance/Quality Control (QA/QC) procedures into their laboratory design and standard operating procedures. LENL-P is certified in several states. In order to perform analyses for AFCEE projects, LENL-P has undergone and complied fully with audits conducted under the auspices of the Air Force. LENL-P's key personnel and their positions responsibilities are outlined in Appendix B-1.
- 1.3.1.4 <u>Geochemical Survey</u> Transglobal Environmental Geochemistry, Texas, (TEG) specializing in ground-water screening, will conduct the geochemical survey. They will perform on site sampling and analysis of the ground water generated during the ground-water screen. TEG's system has been recognized and approved by the Texas Water Commission (TWC) as a viable tool for studying ground-water contaminated plumes.
- 1.3.1.5 <u>Surveying Subcontractor</u> A local engineering and surveying firm, certified in the state of Texas, will be 3517-0111.10

subcontracted to survey boring/sampling locations and elevations and prepare site maps. The firm will report directly to the Site Manager during their survey activities and to the Project Manager prior to and after field work. Their in-house Professional Engineer (PE)/Registered Land Surveyor (RLS) will perform the field surveying and preparation of the final site map. These individuals will follow the AFCEE guidance documents, provided by Law, to prepare their draft and final site map.

1.3.1.6 <u>Ensite - Environmental Contractor - Ensite will be</u> contracted to perform services related to handling, transport and disposal of investigative derived wastes generated during the field investigation. Ensite, located in Atlanta, Georgia, is a whollyowned subsidiary of Law Environmental and is a licensed site remediation contractor.

1.3.2 <u>Key Project Individuals</u>

Key project participants for this project include the project principal, project manager, project chemist, project geologist/site manager, health and safety officer, laboratory manager, and field work parties. The following paragraphs provide a description of the proposed project assignments and responsibilities, a list of individuals expected to serve in each capacity, and a brief synopsis of the participants' related experience.

1.3.2.1 <u>Project Principal</u> - The Project Principal provides technical quality control, oversight and direction for all aspects of the site investigation and data evaluation. Mr. E. Fred Sharpe, Jr., P.E., will serve as the Project Principal. Mr. Sharpe, a Principal with Law Environmental, will serve as the senior reviewer of Law's reports of the Carswell AFB RFI sites. Mr. Sharpe has over 24 years experience with Law Companies.

- 1.3.2.2 <u>Project Manager</u> The Project Manager is responsible for the overall management of the Carswell AFB RFI. He coordinates between office and field personnel, manages administrative requirements, and supervises schedules, technical approach, implementation, and report preparation. Mr. John O'Brien will serve as project manager. Mr. O'Brien has six years of professional experience in site and project management. He is presently managing projects under RCRA, CERCLA, and IRP guidance including Carswell AFB.
- 1.3.2.3 <u>Site Manager</u> The Site Manager will coordinate and supervise the field investigation activities. The Site Manager may also serve as the Health and Safety Officer for the field operations, and will be responsible for the implementation of the Work Plans and Health and Safety Plans. The Site Manager's primary responsibility is the health and safety of the workers on site. Mr. Tom McComb will serve as the Site Manager for this project. Mr. McComb is a Professional Geologist with over seven years experience in site investigations and assessments.
- 1.3.2.4 <u>Project Chemist</u> The Project Chemist is responsible for preparing and implementing the field sampling, preservation, chain-of-custody, and shipping activities. The Project Chemist also performs data evaluation on the chemical data. Ms. Darice Kurtzer will serve as the Project Chemist for this investigation. Ms. Kurtzer is a chemist with over three years of experience in environmental chemistry, Quality Assurance/Quality Control (QA/QC) and data evaluation.
- 1.3.2.5 <u>Laboratory Project Manager</u> The Laboratory Project Manager is responsible for the handling and analysis of water and/or soil samples received by the laboratory. This person oversees sample travel through the lab, analytical procedures,

quality control, reporting and sample disposal. Mr. James Tucci is LENL's Chemical Laboratory Manager. His areas of expertise include environmental field studies, laboratory analyses and personnel management.

- 1.3.2.6 <u>Work Party</u> The work party performs on-site tasks contained in this plan, including drilling, monitoring well installation, and analytical sampling under the direction of the Site Manager.
- 1.3.2.7 Health and Safety Officer The Health and Safety Officer oversees the Health and Safety Plan for Carswell AFB. and Safety Officer conducts personnel training, administers company hazard assessment and surveillance medical program, and coordinates with the Site Manager for site safety. Dr. Jack Peng is Law Environmental's Environmental Health and Safety Officer and is available for consultation during the actual investigation. He is a Certified Industrial Hygienist with over ten years experience in hazardous waste site investigations. Subcontractors working on the Carswell AFB sites under this project will receive a copy of the project Health and Safety Plan. The Law Environmental Site Manager or an assigned qualified Health and Safety Officer will review with subcontractor personnel the Health and Safety Plan prior to any field activities. Site personnel will be required to sign and date the master field logbook to ensure that they understand the safety regulations and procedures as outlined in the Health and Safety Plans.
- 1.3.2.8 <u>Project Safety Officer</u> The Project Safety Officer is responsible for the project Site Safety and Health Plan requirements. Ms. Cindy Kahout will serve as the project Safety Officer for this investigation at Carswell AFB. Ms. Kahout is an Industrial Hygienist with nine years of experience in developing and auditing health and safety programs.

Law Environmental, Inc., has established a strong internal QA program with an associated QA Manual, Engineering Procedures Manual, Equipment Calibration Procedures Manual, and specialty manuals for hazardous waste site investigations and software documentation. Law Environmental employees use these manuals as the basis for conducting all company work within the QA program.

Mr. Leonard Ledbetter, the President of Law Environmental, Inc., has overall responsibility for Law's Corporate QA. The Corporate QA Office is directed by Dr. James R. Wallace, who is responsible for daily management and auditing of Law Environmental's QA Program. The Law Environmental lines of QA responsibility and audit flow from Corporate QA (Mr. Ledbetter and Dr. Wallace) to the Project Principal (Mr. E. Fred Sharpe). This line of QA is outside of the operational lines of authority for this project.

Our Quality Assurance Program is one of the most important factors contributing to client satisfaction and our continued success. The purpose of the program is to provide the client with confidence that services are performed correctly.

Our Principal Review System is an integral part of the Quality Assurance Program. Each project is assigned to a Principal Reviewer who is responsible for maintaining the required professional quality from beginning to completion of the project. Every proposal and report must be reviewed and signed by two people and at least one must be a Law Principal with credentials and experience relevant to the area of work.

In each office, the Chief Scientist or Engineer is responsible for operation of the Quality Assurance Program and for reviewing the performance of assigned Principals. The Chief Engineer conducts internal audits of projects and at least once each year a formal

corporate audit is conducted by Chief Engineers and Principals from other offices. In addition to these internal and corporate audits, the company participates in laboratory and peer review programs sponsored by organizations such as the National Bureau of Standards, U.S. Geological Survey, USEPA, AFCEE and the U.S. Army Corps of Engineers.

Law Companies encourages staff membership and participation in various relevant professional societies. We believe such involvement fosters individual professional growth and helps keep our staff up-to-date on current developments. Many Law Principals also serve on subcommittees or teach in sponsored workshops.

Law will control the quality of subcontractor furnished data and services by source evaluation and selection, evaluation objective evidence of quality compliance to procurement documents furnished by the subcontractor, site and/or source inspections, audits, and/or examination of items or services upon delivery or completion. Prior to award of subcontract, potential suppliers of quality affecting items or services will be evaluated by Law The evaluation may be performed by any appropriate means, including but not limited to, surveys, inspections, audits, or surveillances, depending upon the complexity of the item or service being supplied. When the Project Principal determines it necessary, on-site audits of subcontractor facilities may be conducted under a quality assurance program. While responsibility for quality remains with the subcontractor under his quality control program, Law will conduct certain quality assurance activities to determine the subcontractors compliance with their own quality program.

1.4 QUALITY ASSURANCE OBJECTIVES FOR MEASUREMENT DATA

Quality Assurance (QA) objectives for measurement data are expressed in terms of precision, accuracy, representativeness,

completeness, and comparability (PARCC). QA objectives provide the mechanism for ongoing control and evaluation of measurement data quality throughout the project and will ultimately be used to define data quality for the various measurement parameters. The achievement of these QA/QC objectives will be accomplished through the collection and analysis of field duplicates, rinsates, trip blanks, and field blanks. The analytical data will be assessed by the internal QC such as method blanks, surrogate recoveries, adherence to holding times, and matrix spike/matrix spike duplicate (MS/MSD) data.

1.4.1 Definitions of QA Criteria

The following sections define the precision, accuracy, representativeness, completeness, and comparability (PARCC) parameters.

1.4.1.1 <u>Precision</u> - "Precision" refers to the reproducibility or degree of agreement among duplicate measurements of a single analyte. It is defined by the USEPA as "a measure of mutual agreement among individual measurements of the same property, usually under prescribed similar conditions. Precision is best expressed in terms of the relative percent difference (RPD). Various measures of precision exist depending upon the prescribed similar conditions" (USEPA, 1980). Analytical precision for a single analyte is expressed as a percentage of the difference between results of duplicate samples and matrix spike duplicates for a given analyte. Relative percent difference is calculated as shown in Section 1.10.2.

Precision will be determined through the collection of duplicates and matrix spikes and matrix spike duplicates for the analytical work performed at Carswell AFB. The sampling chemist will select

1 sample in 20 (or one per batch, whichever is fewer) for that matrix and split the sample into three aliquots. The first aliquot will be analyzed routinely for the parameters of interest, while the other two aliquots will be spiked with known quantities of the parameters of interest prior to analysis. The relative percent difference (RPD) will be calculated and used as an indication of the precision for the analyses performed.

During the collection of data using field methods and/or instrumentation, precision is checked by reporting several measurements taken at one location and comparing the results. Precision will be reported as the RPD for two results and as the standard deviation for three or more results. Sample collection precision will be measured in the laboratory with the analysis of field replicates and laboratory duplicates.

1.4.1.2 Accuracy - "Accuracy" refers to the degree of difference between measured or calculated values and the true value. defined by the USEPA as "the degree of agreement of a measurement (or an average of measurements of the same thing), X, with an accepted reference or true value, T, usually expressed as the difference between the two values, X - T, or the difference as a percentage of the reference or true value, 100 (X-T)/T, and sometimes expressed as a ratio, X/T. Accuracy is a measure of bias in a system" (USEPA, 1980). The closer the numerical value of the measurement comes to the true value, or actual concentration, the more accurate the measurement. Analytical accuracy may be expressed as the percent recovery of an analyte which has been added to the environmental sample at a known concentration before analysis. For example, accuracy can be determined from the results of matrix spike analyses performed at the rate of 1 set every 20 samples or 1 per batch. The equation used to calculate percent recovery can be found in Section 1.13.1. The accuracy of simple, yet fundamental, field analyses is difficult to assess quantitatively. Sampling accuracy can be maximized, however, by the adoption and adherence to a strict QA program. Specifically, all procedures will be documented as standard protocol and all equipment and instrumentation will be properly calibrated and well maintained. Trip blanks and equipment rinsates will be included in all sample batches to ensure all samples represent the particular site from which they were taken and to assess the potential for any cross-contamination that may have occurred. In addition to equipment operation and standard operating procedures, a high level of accuracy will be maintained by thorough and frequent review of field procedures. In this manner, any deficiencies will be quickly documented and corrected.

- 1.4.1.3 Representativeness "Representativeness" indicates the degree to which a sample contains the characteristics of the whole from which it came. As defined by the USEPA, representativeness "expresses the degree to which data accurately and precisely represent a characteristic of a population, parameter variations at a sampling point, a process condition, or an environmental condition" (USEPA, 1980). Only qualitative goals for representativeness can be set.
- 1.4.1.4 <u>Completeness</u> "Completeness" is a measure of the amount of data collected with respect to project requirements. It is defined by the USEPA as "a measure of the amount of valid data obtained from a measurement system compared to the amount that was expected to be obtained under correct normal conditions" (USEPA, 1980). Completeness refers to the project as a whole.
- 1.4.1.5 <u>Comparability</u> "Comparability" is a measure of the consistency of the measurement process. It is defined by the USEPA as the parameter that "expresses the confidence with which one data set can be compared to another" (USEPA, 1980). Comparability is

also only expressed in a qualitative manner. To ensure comparability, only established methods and protocols will be used to collect and analyze samples.

1.4.1.6 <u>Analytical Levels</u> - Different levels of quality are required for the analytical data depending upon the planned use of the data. The procedures and methods will vary depending on the level of analytical data required. The USEPA has defined five analytical levels (USEPA, 1987a); these levels are presented in Table 1-3.

1.4.2 Goals

Our goals for this project are designed to be realistic, achievable and ensure field and analytical data quality. Method-specific goals for precision and accuracy are presented in Section 1.10.3. Representativeness and comparability are qualitative goals. Soil representativeness will be maximized through the thorough mixing of the sample, during collection and agreement of best location before sample collection.

Comparability goals will be established by adhering to the use of standard, published sampling and analytical methods and the use of QC samples. The analytical methods completeness goals are 90 percent. However, parameter specific completeness goals for the soil samples collected at Carswell AFB during the RFI will not be stated because matrix interferences and their impact on these goals are unknown at this time. The overall completeness goal (valid data) for Carswell AFB data will be 90 percent. Failure to meet these goals may result in qualification of the data, non-use of the data, or resampling.

TABLE 1-3

ANALYTICAL LEVELS RCRA Facility Investigation Carswell Air Force Base, Texas

ANALYTICAL LEVEL	EL DATA USES	DATAQUALITY	LIMITATIONS	FIELD ACTIVITIES
Level	Site characterizationMonitoring during implementation	 If instruments calibrated and data interpreted correctly, can provide indication of contamination 	 Instruments respond to naturally—occurring compounds 	 HNu Explosimeter Draeger tubes Geophysical surver Geochemical surver
Level II	 Site characterization Evaluation of alternatives Engineering design Monitoring during implementation 	 Dependent on QA/QC steps employed Data typically reported in concentration ranges 	 Tentative ID Techniques/instruments limited mostly to volatiles, metals 	- Temperature - Turbidity - Water level - pH - Specific conductance
Level III	 Risk assessment PRP determination Site characterization Evaluation of alternatives Engineering design Monitoring during implementation 	 Similar detection limits to CLP Less rigorous QA/QC 	Tentative ID in some casesCan provide data of same quality as Level IV	 Ground-water samples Soil samples: surface Soil samples: subsurface Surface water samples Sediment samples Containerized Waste Material
Level IV			Tentative Identificationof non-HSL parametersSome time may be requiredfor validation of packages	Not Applicable
Level V			 May require method development/modification Mechanism to obtain services requires special lead time 	Not Applicable
SOURCE: USEPA, 1987a	987a			187

In establishing the goals for precision and accuracy for each parameter and matrix, the following were considered:

Precision

- · Precision of instrument calibration
- · Precision of duplicate spikes
- · Precision of field duplicates
- Sampling and handling procedures

Accuracy

- · Proper instrument set-up
- Accuracy and stability of the calibration
- · Recovery of spiked samples
- · Recovery of spiked blanks and/or control samples
- Method contamination
- · Surrogate recoveries (if applicable)
- · Interferences
- · Field contamination
- Sampling and handling procedures

The worst performer of all the QC processes for all analytes within each parameter was used to determine the first estimate of the overall goal for each parameter and matrix. In order to provide a realistic approach in case there are unanticipated matrix effects, this estimate of the goal was then relaxed slightly in order to set goals that are realistic and achievable. The overall goal is to allow the use of as much reliable data as possible during this investigation. Data for individual analytes for which any applicable QC result falls outside of the acceptance limits set by the method will be qualified.

Representativeness and comparability are qualitative goals. Representativeness will be dependent upon the careful selection of

sampling locations. Comparability will be established by adhering to the use of standard, published sampling and analytical methods and reporting data in standard units.

1.5 <u>SAMPLING PROCEDURES</u>

Sampling procedures consist of sample collection, transport and storage protocols. These protocols which pertain to the field activities are presented in detail in the Field Sampling Plan (FSP) and are only referenced here.

1.5.1 Sampling Protocols

Sample collection, transport and storage guidance documents to be adhered to are listed below:

- Practical Guide for Ground-Water Sampling, EPA 600/2-85/104, September 1985 (USEPA, 1985)
- Test Methods for Evaluating Solid Wastes, Third Edition, EPA SW-846, November 1986 (USEPA, 1986)
- Handbook to Support the Installation Restoration Program (IRP)
 Statements of Work, May 1991 (AFCEE, 1991)

Detailed sampling protocols are presented in Section 2.0.

1.5.2 Sample Handling

The sample containers, sample volume, method of preservation, shipping and handling procedures are presented in Section 2.2.2.

1.6 SAMPLE CUSTODY

Sample custody procedures during the collection of samples in the field and sample receipt in the laboratory are discussed in the following subsections.

1.6.1 Field Operations

Maintenance of sample custody in the field starts with the collection, preservation and labeling of the sample. Documentation of field sampling activities will be performed. Specific procedures for the maintenance and documentation of the custody of the samples is discussed in detail in Section 2.2.3.

1.6.2 <u>Laboratory Operations</u>

Sample custody, handling, and identification in the laboratory is presented in Appendix B.

1.7 FIELD EQUIPMENT CALIBRATION PROCEDURES

Several instruments will be used in the field investigation. These instruments consist of a photoionization detector (HNu), pH meter, combination temperature and specific conductance meter, and a turbidimeter. Each field meter will be calibrated before use. The calibration will be documented in the field book. Details of calibration and maintenance are presented in Section 2.3 of the FSP. The equipment manuals are presented in Appendices C, D, E and F.

The following sections identify the analytical methods to be utilized.

1.8.1 <u>Identification of Methods</u>

The analytical methods to be utilized are presented in Tables 1-4 (soil, sediment) and 1-5 (ground water, surface water).

1.8.2 Detection Limits and Quantitation Criteria

1.8.2.1 <u>Terminology</u> - Each analytical parameter concentration will be reported as a specific number or less than the Instrument Detection Limit for inorganic methods and less than the Practical Quantification Limit for organic methods. Water values will be reported in parts per billion (ppb, μ g/L) concentrations for organics and in parts per million (ppm, mg/L) for inorganics. Soil values will be reported in parts per million (ppm, mg/kg) for all analyses. Method Detection Limits (MDLs) are defined below.

1.8.2.2 <u>Procedures</u> - The following procedures are used to establish limits of detection and quantification.

Method Detection Limits (MDL) limits are established using the required EPA procedure specified in 40 CFR Part 136 Appendix B. The MDL is the minimum concentration of a substance that can be measured and reported with 99 percent confidence that the value is above zero. A data pool of at least seven standards analyzed at a concentration approximately three times the anticipated MDL is generated. The MDL is estimated by employing the "t" distribution with a 99 percent confidence interval by the following equation:

ANALYTICAL REQUIREMENTS - SOIL AND SEDIMENT SAMPLES RCRA Facility Investigation Carswell Air Force Base, Texas

<u> </u>		
		HOLDING
ANALYTICAL PARAMETER (a)	METHOD (b)	TIME (c)
Volatile Organics	8240	A: 14d
Aromatic Volatile Organics	8020	A: 14d
Total Recoverable Petroleum Hydrocarbons	418.1	P: 14d A: 40d
Metals, total:		A: 6m (unless noted)
1. Aluminum	3050/6010	
2. Antimony	3050/6010	
3. Arsenic	3050/7060 (d)	
4. Barium	3050/6010	
5. Beryllium	3050/6010	
6. Cadmium	3050/6010	
7. Calcium	3050/6010	
8. Chromium	3050/6010	
9. Cobalt	3050/6010	
10. Copper	3050/6010	
11. Iron	3050/6010	
12. Lead	3050/7421 (d)	
13. Mercury	7471	A: 28d
14. Magnesium	3050/6010	
15. Manganese	3050/6010	
16. Molybdenum	3050/6010	
17. Nickel	3050/6010	
18. Potassium	3050/6010	
19. Selenium	3050/7740 (d)	
20. Silver	3050/6010	
21. Sodium	3050/6010	
22. Thallium	3050/7841 (d)	
23. Vanadium	3050/6010	
24. Zinc	3050/6010	

TABLE 1-4

ANALYTICAL REQUIREMENTS - SOIL AND SEDIMENT SAMPLES RCRA Facility Investigation Carswell Air Force Base, Texas

187 51

ANALYTICAL PARAMETER (a)	METHOD (b)	HOLDING TIME (c)
TCLP	1311	See Below

	From: Field Collection to: TCLP extraction	From: TCLP extraction to: Preparative extraction	From: Preparative extraction to: determinative analysis	Total elapsed time
Volatiles	14d	NA	14d	28d
Semi-Volatiles	14d	7d	40d	61d
Mercury	28d	NA	28d	56d
Metals, except Mercury	180d	NA	180d	360d

NOTES:

- (a) For list of analytes and reporting limits, see Table 1-6.
- (b) <u>Test Methods for Evaluating Solid Waste</u> SW-846, 3rd Edition, November 1986 (unless otherwise noted)
- (c) holding time from date of sampling to preparation (P) or analysis (A) $d = days \qquad m = months \qquad h = hours$
- (d) Screening results by method 3050/6010 are acceptable if sample result is greater than 10 times the method detection limit (before correction to dry weight)

TABLE 1-5

ANALYTICAL REQUIREMENTS - GROUND AND SURFACE WATER SAMPLES
RCRA Facility Investigation
Carswell Air Force Base, Texas

ANALYTICAL PARAMETER (a)	METHOD (b)	HOLDING TIME (c)
Volatile Organics	8240	A: 14d
Aromatic Volatile Organics	8020	A: 14d
Extractable Organics	3010 or 3020/8270	P: 7d A: 40d (d)
Petroleum Hydrocarbons	8015 M	A: 14d
Total Recoverable Petroleum Hydrocarbons	418.1	A: 28d (d)
Oil and Grease	9071	A: 28d
Purgeable Halocarbons	5030/8010	A: 14d
Metals, total: 1. Aluminum 2. Antimony 3. Arsenic 4. Barium 5. Beryllium 6. Cadmium 7. Calcium 8. Chromium 9. Cobalt 10. Copper 11. Iron 12. Lead 13. Mercury 14. Magnesium 15. Manganese 16. Molybdenum 17. Nickel 18. Potassium 19. Selenium 20. Silver 21. Sodium 22. Thallium 23. Vanadium 24. Zinc	3005/6010 3005/6010 7060 (e) 3005/6010	A: 6m (unless noted) A: 28d

TABLE 1-5

ANALYTICAL REQUIREMENTS - GROUND AND SURFACE WATER SAMPLES RCRA Facility Investigation Carswell Air Force Base, Texas

ANALYTICAL PARAMETER (a)	METHOD (b)	HOLDING TIME (c)
TCLP	1311	See Below

	From: Field Collection to: TCLP extraction	From: TCLP extraction to: Preparative extraction	From: Preparative extraction to: determinative analysis	Total elapsed time
Volatiles	14d	NA	14d	28d
Semi-Volatiles	14d	7d	40d	61d
Mercury	28d	NA	28d	56 d
Metals, except Mercury	180d	NA	180d	360d

NOTES:

- (a) For list of analytes and reporting limits, see Table 1-7.
- (b) <u>Test Methods for Evaluating Solid Waste</u> SW-846, 3rd Edition, November 1986 (unless otherwise noted)
- (c) holding time from date of sampling to preparation (P) or analysis (A) d = days m = months h = hours
- (d) Extracts to be analyzed within 40 days of date of extraction.
- (e) Screening results by method 3005/6010 are acceptable if sample result is greater than 10 times the method detection limit (before correction to dry weight)

$$MDL = (t)(S)$$

where:

S = is the standard deviation of the data pool

Instrument Detection Limits (IDL) are established by the EPA CLP protocol, which allows for generating a data pool by analyzing a minimum of seven standards at three times the anticipated IDL on three non-consecutive days. The standard deviation (S) of the seven data for each of the three days is calculated and the IDL established by the following equation:

$$IDL = \frac{S1 + S2 + S3}{3} * 3$$

The Practical Quantitation Limit (PQL) is the lowest level that can be reliably achieved within specified limits of precision and accuracy during routine laboratory operating conditions. PQL is calculated as follows:

$$PQL = 10 * S$$

where:

S = is the standard deviation of the data used to determine MDL

The matrix used for these studies is spiked reagent water processed through the appropriate analytical procedure.

The Maximum Quantitation Limit (MQL) is that limit established by AFCEE as being the maximum allowable quantitation limit of an analyte. These limits are based upon the results of spiking at the estimated quantitation limit in accordance with EPA SW-846 protocols, or the results of matrix spiking of past HSD/YAQ samples.

The Maximum Quantitation Limit (MQL) for each parameter is calculated as follows:

MQL = F * S

where:

S = is the standard deviation of the data used to determine MDL

F = is a factor between five and ten based on chromatographic behavior

The matrix used for these studies is spiked reagent water processed through the appropriate analytical procedure.

1.8.2.3 <u>Values</u> - The reporting limits required for this project are presented in Tables 1-6 and 1-7 for soil and sediment and ground water and surface water. The laboratory established detection and quantitation limits are presented in Appendix A.

1.8.3 Method Calibration

The calibration procedures, preparation of calibration standards, and frequency of initial and continuing calibration checks are described for each analytical method in the following subsections.

TABLE 1-6

PROJECT-REQUIRED REPORTING LIMITS - SOIL/SEDIMENT Remedial Investigation/Feasibility Study Carswell Air Force Base, Texas

ANALYTICAL				REPOR	
METHOD		POUND	UNITS	IDL	MQL
6010 (SW846)	Aluminum		mg/kg	10	50
,	Antimony		mg/kg	5.6	15
	Arsenic		mg/kg	4.2	30
	Barium		mg/kg	3	10
	Beryllium		mg/kg	0.2	1
	Cadmium		mg/kg	0.4	0.5
	Calcium		mg/kg	10.2	100
	Chromium		mg/kg	8.0	5
	Cobalt		mg/kg	1.3	5
	Copper		mg/kg	8.0	5
	Iron		mg/kg	5	5
	Lead		mg/kg	3.7	5
	Magnesium		mg/kg	11.7	100
	Manganese		mg/kg	0.6	2
	Molybdenum		mg/kg	3.6	10
	Nickel		mg/kg	2.5	15
	Potassium		mg/kg	13.4	100
	Selenium		mg/kg	7.9	50
	Silver		mg/kg	8.0	5
	Sodium		mg/kg	25.5	100
	Thallium		mg/kg	7	7
	Vanadium		mg/kg	1	10
	Zinc		mg/kg	0.7	2
7060 (SW846)	Arsenic		mg/kg	0.2	0.5
7421 (SW846)	Lead		mg/kg	0.1	0.5
7740 (SW846)	Selenium		mg/kg	0.1	0.5
7841 (SW846)	Thallium		mg/kg	0.1	7
7471 (SW846)	Mercury		mg/kg	0.02	0.1
1311/7060	Arsenic		mg/kg	0.2	0.5
1311/6010	Barium		mg/kg_	3	10
1311/6010	Cadmium		mg/kg	0.4	0.5
1311/6010	Chromium		mg/kg	0.8	5_
1311/7421	Lead		mg/kg	0.1	0.5_
1311/7471	Mercury		mg/kg	0.02	0.1
1311/7740	Selenium		mg/kg	0.1	0.5
1311/6010	Silver		mg/kg	0.8	5

PROJECT-REQUIRED REPORTING LIMITS - SOIL/SEDIMENT Remedial Investigation/Feasibility Study Carswell Air Force Base, Texas

ANALYTICAL			REPOR LIM	
METHOD	COMPOUND	UNITS	PQL	MQL
8240 (SW846)	Acetone	mg/kg	0.001	0.01
0240 (344040)	Benzene	mg/kg	0.001	0.005
	Bromodichloromethane	mg/kg	0.001	0.005
	Bromoform	mg/kg	0.005	0.005
	Bromomethane	mg/kg	0.001	0.01
	2-Butanone (MEK)	mg/kg	0.01	0.01
	Carbon disulfide	mg/kg	0.005	0.005
	Carbon tetrachloride	mg/kg	0.005	0.005
	Chlorobenzene	mg/kg	0.005	0.005
	Chloroethane	mg/kg	0.005	0.01
	2—Chloroethyl vinyl ether	mg/kg	0.01	0.01
	Chloroform	mg/kg	0.005	0.005
	Chloromethane	mg/kg	0.001	0.01
	Dibromochloromethane	mg/kg	0.005	0.005
	1,1-Dichloroethane	mg/kg	0.005	0.005
	1,2-Dichloroethane	mg/kg	0.005	0.005
	1,1-Dichloroethene	mg/kg	0.005	0.005
	trans-1,2-Dichloroethene	mg/kg	0.005	0.005
	1,2-Dichloropropane	mg/kg	0.001	0.005
	cis-1,3-Dichloropropene	mg/kg	0.001	0.005
	trans-1,3-Dichloropropene	mg/kg	0.005	0.005
	Ethylbenzene	mg/kg	0.005	0.005
	2-Hexanone	mg/kg	0.005	0.01
	Methylene chloride	mg/kg	0.005	0.005
	4-Methyl-2-pentanone (MIBK)	mg/kg	0.01	0.01
	Styrene	mg/kg	0.005	0.005
	1,1,2,2—Tetrachloroethane	mg/kg	0.005	0.005
	Tetrachloroethene	mg/kg	0.005	0.005
	Toluene	mg/kg	0.005	0.005
	1,1,1-Trichloroethane	mg/kg	0.005	0.005
	1,1,2-Trichloroethane	mg/kg	0.005	0.005
	Trichloroethene	mg/kg	0.005	0.005
	Vinyl acetate	mg/kg	0.01	0.01
	Vinyl acetate Vinyl chloride	mg/kg	0.002	0.01
	Xylenes (total)	mg/kg	0.005	0.005
	1,2-Dichloroethane-d4 (surrogate)	mg/kg	NA	NA
	Toluene-d8 (surrogate)	mg/kg	NA NA	NA
	4-Bromofluorobenzene (surrogate)	mg/kg	NA NA	NA NA

PROJECT-REQUIRED REPORTING LIMITS - SOIL/SEDIMENT Remedial Investigation/Feasibility Study Carswell Air Force Base, Texas

ANALYTICAL			REPOR'	
METHOD	COMPOUND	UNITS	PQL	MQL
8020 (SW846)	Benzene	mg/kg	0.0005	0.001
0020 (011040)	Toluene	mg/kg	0.0005	0.001
	Ethylbenzene	mg/kg	0.0005	0.002
	Xylenes	mg/kg	0.003	0.004
	Fluorobenzene (surrogate)	mg/kg	NA	NA
	4-Chlorotoluene (surrogate)	mg/kg	NA NA	NA
9070/418.1 (EPA600)	Total Recoverable Petroleum Hydrocarbons	mg/kg	10	30
418.1	Total Recoverable Petroleum Hydrocarbons	mg/kg	10	30
1311/8240	Benzene	mg/kg	0.001	0.005
	Carbon tetrachloride	mg/kg	0.005	0.005
	Chlorobenzene	mg/kg	0.005	0.005
	Chloroform	mg/kg	0.005	0.005
	1,2-Dichloroethane	mg/kg	0.005	0.005
	2-Butanone (MEK)	mg/kg	0.01	0.01
	Vinyl Chloride	mg/kg	0.002	0.01
1311/8270	Pentachlorophenol	mg/kg	0.33	1
	2,4,5-Trichlorophenol	mg/kg	0.033	1.6
	2,4,6-Trichlorophenol	mg/kg	0.033	0.3
	1,4-Dichlorobenzene	mg/kg	0.033	0.3
	2,4-Dinitrotoluene	mg/kg	0.033	0.3
	Hexachlorobenzene	mg/kg	0.033	0.3
	Hexachlorobutadiene	mg/kg	0.033	0.3
	Hexachloroethane	mg/kg	0.033	0.3
	Nitrobenzene	mg/kg	0.033	0.3

NA - Not applicable.

NE - Not established at this time.

IDL - Instrument Detection Limit

MQL - Maximum Quantitation Limit

ANALYTICAL	COMPOUND	UNITS	REPORTI	NG LIMIT
METHOD			IDL	MQL
6010 (SW846)	Aluminum	mg/L	0.1	0.5
	Antimony	mg/L	0.056	0.5
	Arsenic	mg/L	0.042	0.3
	Barium	mg/L	0.03	0.1
	Beryllium	mg/L	0.002	0.01
	Cadmium	mg/L	0.004	0.005
	Calcium	mg/L	0.102	1
	Chromium	mg/L	0.008	0.05
	Cobalt	mg/L	0.013	0.05
	Copper	mg/L	0.008	0.05
	Iron	mg/L	0.05	0.05
	Lead	mg/L	0.037	0.2
	Magnesium	mg/L	0.117	1
	Manganese	mg/L	0.006	0.02
	Molybdenum	mg/L	0.036	0.1
	Nickel	mg/L	0.025	0.15
	Potassium	mg/L	0.134	5
	Selenium	mg/L	0.079	1
	Silver	mg/L	0.008	0.05
	Sodium	mg/L	0.255	1
	Thallium	mg/L	0.072	0.4
	Vanadium	mg/L	0.01	0.1
	Zinc	mg/L	0.007	0.02
7060 (SW846)	Arsenic	mg/L	0.002	0.005
7421 (SW846)	Lead	mg/L	0.001	0.005
7740 (SW846)	Selenium	mg/L	0.001	0.005
7471 (SW846)	Mercury	mg/L	0.0002	0.001

TABLE 1-7

			REPORTING LIMIT		
ANALYTICAL METHOD	COMPOUND	UNITS	MDL	PQL	MQL
8240 (SW846)	Acetone		7.5	10	10
	Benzene	µg/L	0.4	1	5
	Bromodichloromethane	µg/L	0.3	1	5
	Bromoform	μg/L	0.2	5	5
	Bromomethane	µg/L	0.7	1	10
	2-Butanone (MEK)	µg/L	2.8	10	10
	Carbon disulfide	µg/L	0.7	5	5
	Carbon tetrachloride	μ g/L	0.4	5	5
	Chlorobenzene	µg/L	0.2	5	5
	Chloroethane	µg/L	0.3	5	10
	2-Chloroethyl vinyl ether	μg/L	8	10	10
	Chloroform	μg/L	0.3	5	5
	Chloromethane	μg/L	0.6	1	10
	Dibromochloromethane	μg/L	0.3	5	5
	1,1-Dichloroethane	μg/L	0.3	5	5
	1,2-Dichloroethane	μ g/L	0.4	5	5
	1,1-Dichloroethene	μg/L	0.3	5	5
	trans-1,2-Dichloroethene	μg/L	0.4	5	5
	1,2-Dichloropropane	μg/L	0.6	1	5
	cis-1,3-Dichloropropene	µg/L	0.6	1	5
	trans-1,3-Dichloropropene	µg/L	0.2	5	5
	Ethylbenzene	μg/L	0.2	5	5
	2-Hexanone	<i>μ</i> g/L	2.1	5	10
	Methylene chloride	µg/L	4.5	5	5
	4-Methyl-2-pentanone (MIBK)	µg/L	7	10	10
	Styrene	μ g/L	0.5	5	5
	1,1,2,2-Tetrachloroethane	μ g/L	0.7	5	5
	Tetrachloroethene	μ g/L	0.2	5	5
	Toluene	μg/L	0.3	5	5

TABLE 1-7

			REPORTING LIMIT		
ANALYTICAL METHOD	COMPOUND	UNITS	MDL	PQL	MQL
	1,1,1-Trichloroethane	μg/L	0.4	5	5
	1,1,2-Trichloroethane	μg/L	0.7	5	5
	Trichloroethene	μg/L	0.2	5	5
	Vinyl acetate	μ g/L	0.7	10	10
	Vinyl chloride	μ g/L	0.6	2	10
	Xylenes (total)	μ g/L	0.7	5	5
	1,2-Dichloroethane-d4 (surrogate)	µg/L	NA	NA	NA
	Toluene-d8 (surrogate)	μ g/L	NA	NA	NA
	4-Bromofluorobenzene (surrogate)	<i>μ</i> g/L	NA ——	NA	NA
8020 (SW846)	Benzene	μg/L	0.5	0.5	1
	Toluene	μ g/L	0.5	0.5	2
	Ethylbenzene	μ g/L	0.5	0.5	4
	Xylenes	μ g/L	1	1	2
	Fluorobenzene (surrogate)	μ g/L	NA	NA	NA
	4-Chlorotoluene (surrogate)	μg/L	NA	NA	NA
8015M (SW846)	Diesel	mg/L	0.072	0.2	1
	Gasoline	mg/L	0.074	0.5	1
9071 (SW846)	Oil & Grease	μg/L	NE	NE	NE
418.1	Total Recoverable Petroleum Hydrocarbons	mg/L	0.4	1	1

	COMPOUND	UNITS	REPORTING LIMIT		
ANALYTICAL METHOD			MDL	PQL	MQL
	TCLP	μg/L			
1311/8240	Benzene	$\mu_{ extsf{g}}/ extsf{L}$	0.4	1	5
	Carbon tetrachloride	μg/L	0.4	5	5
	Chlorobenzene	<i>μ</i> g/L	0.2	5	5
	Chloroform	μg/L	0.3	5	5
	1,2-Dichloroethane	μ_{g}/L	0.4	5	5
	2-Butanone (MEK)	<i>μ</i> g/L	2.8	10	10
	Vinyl chloride	<i>μ</i> g/L	0.6	2	10
1311/8270	Pentachlorophenol	μg/L	0.6	10	30
	2,4,5-Trichlorophenol	μg/L	1	1	50
	2,4,6-Trichlorophenol	μg/L	0.7	1	10
	1,4-Dichlorobenzene	μg/L	0.7	1	10
	2,4-Dinitrotoluene	<i>μ</i> g/L	0.8	1	10
	Hexachlorobenzene	<i>μ</i> g/L	0.5	1	100
	Hexachlorobutadiene	<i>µ</i> g/L	0.7	1	10
	Hexachloroethane	μg/L	0.9	1	10
	Nitrobenzene	μg/L	0.9	1	10

TABLE 1-7

			IDL	MQL
1311/6010	Barium	mg/L	0.030	0.1
1311/6010	Cadmium	mg/L	0.004	0.005
1311/6010	Chromium	mg/L	0.008	0.05
1311/7421	Lead	mg/L	0.001	0.005
1311/7471	Mercury	mg/L	0.0002	0.001
1311/7740	Selenium	mg/L	0.001	0.005
1311/6010	Silver	mg/L	0.008	0.05

NA - Not applicable.

NE - Not established at this time.

MDL - Method Detection Limit

MQL - Maximum Quantitation Limit

PQL - Practical Quantitation Limit

^{*} Project - Required Reporting Limits for methodologies performed by TEG are listed in the TEG QA Manual in Appendix B-2.

At a minimum, all instruments and other equipment used by Law will be calibrated and maintained at the recommended intervals prescribed by the analytical method employed. In those cases where it has been demonstrated that more frequent calibration or maintenance is required, the base method will be enhanced as necessary. Calibration checks and preventive maintenance procedures are permanently recorded in a log book for future reference. Whenever possible, calibration checks on instruments and support equipment are performed using reference materials (e.g., weights) which are traceable to the EPA Environmental Monitoring Systems Laboratory (EMSL) or bear certificates of standardization from the National Institute of Standards and Technology.

All chemical standards used by Law are certified to have the highest reasonable purity with assays or certificates of analysis provided with material. All materials are purchased from established and reputable chemical suppliers. All standard materials are dated and stored under appropriate conditions in order to maintain chemical stability. Each time a specific standard is prepared, the material mass and dilution information is recorded in the standards record book. All prepared standards (stock and working) are labeled in reference to the book number, page number, and entry number of the Analytical Standards Preparations Record Book. Additionally, the description, date, and analyst are given on the standard label.

1.8.3.1 Organics by Gas Chromatography (GC) with Second Column Confirmation - Analyses performed on the GC will follow EPA Methods. Second column confirmation is required for samples which exhibit a positive result. The confirmation system must contain a dissimilar column and is calibrated and subject to the same QC as the primary GC system. The primary or secondary result may be reported. The two results will not be averaged.

Prior to calibration, retention time windows for each standard on each GC column are determined whenever a new GC column is installed as follows.

Three injections of each standard are made over a 72-hour period at approximately equal intervals. A standard deviation is calculated from the three absolute retention times. For multi-response analytes, one major peak is chosen from the envelope for the retention time study. Retention time windows for each analyte are updated daily and are equal to the continuing calibration analyte retention time plus or minus three times the standard deviation determined in the study. If the retention time window for an analyte is zero, a standard deviation of a similar compound is used.

Tentative identification of an analyte occurs when a peak from a sample falls within the daily retention time window. Multi-response analytes are identified primarily by pattern recognition.

Calibration procedures for both systems are presented below.

- 1. GC system is checked daily prior to analysis of samples by reviewing temperatures to injector parts, detectors and columns, verifying that the system is method-specific to ensure reliability and reproducibility of analytical results.
- 2. GC is calibrated with five standards and a calibration blank using external standard technique.
- 3. If the %RSD of the RF (internal standard technique) or CF (external standard technique) for the five standards is less than or equal to 20 percent, then the average RF is used for quantitation. If the RF is greater than 20 percent, a calibration curve is established by plotting response versus amount.

- After development of each new five-point calibration and at the start of every analysis day, a mid-point daily initial calibration verification (ICV) containing all single peak analytes of interest is run. Pesticide/PCB standards also include Toxaphene, Chlordane, and Aroclors The standard's RF or CF must agree with the initial calibration average RF or CF within plus or minus 15 percent D. For a calibration curve, the concentration of the mid-point initial calibration standard is compared to the ICV concentration. The percent recovery of the ICV is obtained using the following formula: Value divided by Mid-Point Initial Calibration True Value (calculated from the curve) times 100. The percent recovery of the ICV must be within plus or minus 15 percent.
- 5. After every ten samples within the sequence and at the end of the sequence, a mid-level continuing calibration verification (CCV) is analyzed which must be within plus or minus 15 percent D of the ICV. For a calibration curve, the concentration of the ICV is compared to the concentration of the CCV. The percent recovery of the CCV must be within plus or minus 15 percent.
- 6. The retention time (RT) for all identified analytes in an analytical run must match the RT windows calculated for the analytical standards.
- 7. Second column confirmation is subject to the proceeding acceptance criteria.
- 1.8.3.2 <u>Volatile and Semi-Volatile Organics by Gas Chromatography/Mass Spectrometry (GC/MS)</u> Analyses performed on the GC/MS will follow EPA Methods. This includes the following calibration procedure:

- 1. Instrument calibration will be performed every 12-hour time period. The GC/MS will be tuned to meet ion abundance criteria given in Tables 1-8 and 1-9 for decafluorotriphenylphosphine (DFTPP) or 4-Bromofluorobenzene (BFB).
- 2. Initial calibration will be performed on calibration standards at five concentration levels containing each compound of interest and each surrogate standard.

The relative retention time (RRT) of each compound in each calibration run should agree within 0.06 RRT units. The average relative response factor (RRF) and percent relative standard deviation (%RSD) is calculated for each The RRF for the System Performance Check compound. (SPCC: Chloromethane, Compounds Volatiles: 1,1-Dichloroethane, Bromoform, 1,2,2-Tetrachloroethane, Chlorobenzene. Semi-volatiles: N-nitroso-di-npropylamine, Hexachlorocyclo-pentadiene, 2,4-Dinitrophenol, 4-Nitrophenol) must be greater than or equal to 0.300 (0.250 for Bromoform) for volatiles and must be greater than or equal to 0.050 for semivolatiles. The %RSD for the Calibration Check compounds (CCC: Volatiles - Vinyl chloride, 1,1-Dichloroethene, Chloroform, 1,2-Dichloropropane, Toluene, Ethylbenzene; Semi-volatiles Base/Neutrals -Acenaphthene, Dichlorobenzene, Hexachlorobutadiene, N-Nitroso-di-n-Di-n-octylphthalate, phenylamine, Fluoranthene, Benzo(a)pyrene; Acids - 4-Chloro-3-methylphenol, 2,4-Dichlorophenol, 2-Nitrophenol, Phenol, Pentachlorophenol, 2,4,6-Trichlorophenol) must be less than 30 percent. The %RSD for the other compounds should be less than 30 percent. If not, a notation is made in the sample documentation.

TABLE 1-8

DFTPP KEY IONS AND ABUNDANCE CRITERIA® Carswell Air Force Base, Texas

Mass	Ion Abundance Criteria	
51	30-60% of mass 198	
68	<2% of mass 69	
70	<2% of mass 69	
127	40-60% of mass 198	
197	<1% of mass 198	
198	Base peak, 100% relative abundance	
199	5-9% of mass 198	
275	10-30% of mass 198	
365	>1% of mass 198	
441	Present but less than mass 443	
442	>40% of mass 198	
443	17-23% of mass 442	

^{*}J.W. Eichelberger, L.E. Harris, and W.L. Budde. "Reference Compound to Calibrate Ion Abundance Measurement in Gas Chromatography-Mass Spectrometry," Analytical Chemistry, <u>47</u>, 995 (1975).

TABLE 1-9
BFB KEY IONS AND ABUNDANCE CRITERIA®

MASS	ION ABUNDANCE CRITERIA
50	15-40% of mass 95
75	30-60% of mass 95
95	base peak, 100% relative abundance
96	5-9% of mass 95
173	less than 2% of mass 174
174	less than 50% of mass 95
175	5-9% of mass 174
176	greater than 95% but less than 101% of mass 174
177	5-9% of mass 176

^{*} EPA Method 8240, SW-846, 3rd Edition, November 1986.

- 3. After the preparation of each new initial five-point calibration curve and after tune criteria are met for each 12-hour time period, the initial calibration is verified (using a midpoint calibration standard containing all analytes) prior to running any samples. This initial calibration verification (ICV) must satisfy the above SPCC criteria. In addition, the percent difference (%D) for the CCC must be less than or equal to 25 percent for the volatiles and less than or equal to 30 percent for the semi-volatiles. If these criteria are not met for any single CCC, corrective action must be taken or a new five point calibration must be performed.
- 4. Internal standards for GC/MS volatiles will be added to each sample, standard and reagent blank. The final concentration of each internal standard will be 50 μ g/L. The internal standards are as follows:
 - Bromochloromethane
 - · 1,4-Difluorobenzene
 - Chlorobenzene-d₅

Table 1-10 presents the analytes quantitated by each internal standard.

Internal standards for GC/MS semi-volatiles will be added to each sample, standard, and reagent blank. The final concentration of each internal standard will be 40 nanograms per microliter (ng/ μ L). The internal standards are as follows:

- · 1,4-Dichlorobenzene-d₄
- · Naphthalene-d_g
- · Acenaphthene-d₁₀
- Phenanthrene-d₁₀

TABLE 1-10

VOLATILE INTERNAL STANDARDS WITH CORRESPONDING ANALYTES ASSIGNED FOR QUANTITATION Carswell Air Force Base, Texas

DDOMOCI II ODOMETI IANE	1.4 DIST HODODENZENE	CUI ODODENZENE 45
BROMOCHLOROMETHANE	1,4-DIFLUOROBENZENE	CHLOROBENZENE-d5
Acetone	Benzene	Bromofluorobenzene*
Acrolein	Bromodichloromethane	Chlorobenzene
Acrylonitrile	Bromoform	Ethylbenzene
Bromomethane	2-Butanone	Ethyl methacrylate
Carbon disulfide	Carbon tetrachloride	2-Hexanone
Chloroethane	Chlorodibromomethane	4-Methyl-2-pentanone
Chloroform	2-Chloroethyl vinyl ether	Styrene
Chloromethane	Dibromomethane	1,1,2,2-Tetrachloroethane
Dichlorodifluoromethane	1,4-Dichloro-2-butene	Tetrachloroethene
1,1-Dichloroethane	1,2-Dichloropropane	Toluene
1,2-Dichloroethane	cis-1,3-Dichloropropene	Toluene-d8*
1,2-Dichloroethane-d4*	trans-1,3-Dichloropropene	1,2,3-Trichloropropane
1,1-Dichloroethene	1,1,1-Trichloroethane	Xylene
trans-1,2-Dichloroethene	1,1,2-Trichloroethane	
lodomethane	Trichloroethene	
Methylene chloride	Vinyl acetate	
Trichlorofluoromethane		
Vinyl chloride		

^{*} Surrogate

187 72

- · Chrysene-d₁₂
- · Perylene-d₁₂

Table 1-11 presents the analytes quantitated by each internal standard.

- 1.8.3.3 <u>Metals by Inductively Coupled Plasma (ICP)</u> Metal analyses on the ICP will follow current EPA SW846 method procedures. The instrument must be calibrated daily or once every 24 hours and each time the instrument is set up. This includes the following calibration procedure:
 - 1. Verify instrument is operating satisfactorily by checking automatic gain setting and optical alignment.
 - 2. Calibrate instrument with matrix matched mixed standards at four concentration levels and a blank. The correlation coefficient must be greater than or equal to 0.995.
 - 3. Verify the calibration with a 2nd source Initial Calibration Verification (ICV) standard. The observed result must be plus or minus ten percent of the expected value.
 - 4. Verify the calibration blank. The observed result must be less than three times the detection limit.
 - 5. Verify high-level standard calibration.
 - 6. Every ten samples, a Continuing Calibration Verification (CCV)/Continuing Calibration Blank (CCB) pair is run which must be plus or minus ten percent of expected value and less than three times detection limit, respectively.

TABLE 1-11

SEMI-VOLATILE INTERNAL STANDARDS WITH CORRESPONDING ANALYTES ASSIGNED FOR QUANTITATION Carswell Air Force Base, Texas

1,4-DICHLOROBENZENE-d4	NAPHTHALENE-d8	ACENAPHTHENE-d10
Aniline	Acetophenone	Acenaphthene
Benzyl alcohol	Benzoic acid	Acenaphthylene
Bis(2-chloroethyl)ether	Bis(2-chloroethoxy)methane	1-Chloronaphthalene
Bis(2-chloroisopropyl)ether	4-Chloroaniline	2-Chloronaphthalene
2-Chlorophenol	4-Chloro-3-methylphenol	4-Chlorophenyl phenyl ether
1,2-Dichlorobenzene	2,4-Dichlorophenol	Dibenzofuran
1,3-Dichlorobenzene	2,6-Dichlorophenol	Diethylphthalate
1,4-Dichlorobenzene	alpha,alpha-Dimethyl-phenethylamine	Dimethylphthalate
Ethyl methanesulfonate	2,4-Dimethylphenol	2,4-Dinitrophenol
2-Fluorophenol*	Hexachlorobutadiene	2,4-Dinitrotoluene
Hexachloroethane	Isophorone	2,6-Dinitrotoluene
Methyl methanesulfonate	2-Methylnaphthalene	Fluorene
2-Methylphenol	Naphthalene	2-Fluorobiphenyl*
4-Methylphenoi	Nitrobenzene	Hexachlorocyclopentadiene
N-Nitrosodimethylamine	Nitrobenzene-d8*	1-Naphthylamine
N-Nitroso-di-n-propylamine	2-Nitrophenol	2-Naphthylamine
Phenol	N-Nitroso-di-n-butylamine	2-Nitroaniline
Phenoi-d6*	N-Nitrosopiperidine	3-Nitroaniline
2-Picoline	1,2,4-Trichlorobenzene	4-Nitroaniline
		4-Nitrophenol
		Pentachlorobenzene
		1,2,4,5-Tetrachlorobenzene
		2,3,4,6-Tetrachlorophenol
		2,4,6-Tribromophenol*
		2,4,5-Trichlorophenol
		2,4,6-Trichlorophenol

^{*} Surrogate

TABLE 1-11

SEMI-VOLATILE INTERNAL STANDARDS WITH CORRESPONDING ANALYTES ASSIGNED FOR QUANTITATION Carswell Air Force Base, Texas

PHENANTHRENE-d10	CHRYSENE-d12	PERYLENE-d12
4-Aminobiphenyl	Benzidine	Benzo(b)fluoranthene
Anthracene	Benzo(a)anthracene	Benzo(k)fluoranthene
4-Bromophenyl phenyl ether	Bis(2-ethylhexyl)phthalate	Benzo(g,h,i)perylene
Di-n-butylphthalate	Butylbenzylphthalate	Benzo(a)pyrene
4,6-Dinitro-2-methylphenol	Chrysene	Dibenz(a,j)acridine
Diphenylamine	3,3'-Dichlorobenzidine	Dibenz(a,h)anthracene
1,2-Diphenylhydrazine	p-Dimethylaminoazobenzene	7,12-Dimethylbenz(a)-anthracene
Fluoranthene	Pyrene	Di-n-octylphthalate
Hexachlorobenzene	Terphenyl-d14*	Indeno(1,2,3-cd)pyrene
N-Nitrosodiphenylamine		3-Methylcholanthrene
Pentachlorophenol		
Pentachloronitrobenzene		
Phenacetin		
Phenanthrene		
Pronamide		

^{*} Surrogate

- 7. Check for interferences. An interference check solution must be analyzed at the beginning and at the end of the run (or at least every eight hours); refer to EPA Methods for acceptable limits which are plus or minus 20 percent. The interference check solution contains aluminum, calcium, iron, and magnesium.
- 1.8.3.4 <u>Cold-Vapor Atomic Absorption</u> Mercury is analyzed using cold-vapor atomic absorption (CVAA). An aliquot of sample is acidified and then undergoes a heated, oxidation digestion with potassium permanganate and potassium persulfate. After digestion, a solution of sodium chloride-hydroxylamine sulfate is added to the sample digestate to reduce excess permanganate and remove free chlorine. A reducing agent (stannous chloride) is then added to the solution, resulting in a reduction of the mercury to an elemental state. The elemental mercury is aerated from the solution in a closed system and the mercury vapor content is measured as it passes through a cell positioned in the light path of an atomic absorption spectrophotometer. The calibration procedure is as follows:
 - 1) Optimize instrument setting and alignment by maximizing the energy setting.
 - 2) Align cell minimizing absorbance reading.
 - 3) Calibrate instrument with standards at five concentration levels and a blank. The correlation coefficient must be greater than or equal to 0.995.
 - 4) Verify calibration by running an initial calibration verification standard (ICV) and a calibration blank (ICB). The observed result of the ICV must be within plus or minus 20 percent of the expected value and the

- observed result of the calibration blank must be less than three times the detection limit.
- 5) Analyze a CCV/CCB pair every ten samples. The response must be within 20 percent of the initial response and less than three times the detection limit, respectively.
- 1.8.3.5 Method E 418.1, Total Recoverable Petroleum Hydrocarbons Total recoverable petroleum hydrocarbons (TRPH) are defined by the method used for their determination. The sample is extracted with trichlorofluoroethane, passed through a silica gel adsorption process, and measured by infrared spectrophotometry. This procedure is based upon EPA's Method E 418.1 and SW-846 9073 (Test Methods For Evaluating Solid Waste, Physical/Chemical Methods, SW-846 3rd Edition, Proposed Update II, June 1990) for TRPHs and 413.2 and SW-846 9071A for oil and grease.
- 1.8.3.5.1 Calibration Calibrate the infrared instrument, using the appropriate series of working standards, using a 1 cm cell for water samples and a 5 cm cell for soil samples. It is not necessary to add silica gel to the standards. Determine absorbance directly for each solution at 2930 cm⁻¹. Prepare a calibration plot of absorbance versus milligrams petroleum hydrocarbons per 100 mL solution on a calculator.
- 1.8.3.6 <u>Metals by Graphite Furnace Atomic Absorption (GFAA)</u> Metal analyses performed on the GFAA will follow current EPA SW 846 Methods. Instrument must be calibrated daily or once every 24 hours and each time the instrument is set up. This includes the following calibration procedures:
 - 1. The lamp must be peaked for position and for wavelength (the temperature of the furnace is automatically calibrated at 2600 degrees centigrade).

- 2. After the proper conditions for each element are programmed for furnace operation, distilled water is injected and run as a sample. This is done several times until the instrument response produces a steady base line absorbance.
- 3. Verify instrument is operating satisfactorily by checking the energy output of the lamp and by checking the characteristic mass on the midpoint standard, which must be plus or minus ten percent of the true value for that standard.
- 4. Calibrate instrument with four standards and a calibration blank. The calibration curve must have a correlation coefficient of greater than or equal to 0.995.
- 5. Verify the calibration with a second source Initial Calibration Verification (ICV) standard. The observed result must be within plus or minus ten percent of the expected result.
- 6. Verify the calibration blank. The blank must be less than three times detection limit.
- 7. Verify low level standard calibration.
- 8. Every ten samples, a Continuing Calibration Verification (CCV) and Continuing Calibration Blank (CCB) pair is run and must be plus or minus ten percent of expected value and less than three times the detection limit, respectively.

The following sections describe the reduction, validation, and reporting of data employed after samples are analyzed.

1.9.1 Data Management

Analytical data are collected and processed in accordance with the requirements of the laboratory's protocols. These requirements include sample documentation and data collection. Sample/data flow is outlined in Appendix B for each laboratory.

1.9.2 Data Reduction

Raw data are to be reduced as specified by each method to produce results in the following units:

	SOLID	<u>AOUEOUS</u>
Organic parameters	mg/kg	μ g/L
Inorganic parameters	mg/kg	mg/L

Note that for the organic and inorganic parameters, solid and sediment results will be corrected for the percentage of solids in the sample ("dry weight").

1.9.3 Data Quality Assessment

The parameters of precision, accuracy, representativeness, completeness, and comparability (PARCC) are indicators of data quality (USEPA, 1987a). Establishing goals for these parameters serves to guide the choice of the analytical methodology. It also establishes a strategy for the evaluation of the data once they

have been acquired to determine whether the goals of the project have been met. Upon their receipt from the laboratory, the chemical analysis data will be evaluated by experienced personnel against pre-determined criteria to determine whether their quality meets the requirements of the project. The laboratory quality control (QC) data and the field QC data will be evaluated to objectively ascertain the level of quality of the data. Any issues requiring clarification by the laboratory or the samplers will be identified and pursued at this point. The data quality level will then be compared to that required by the project. If problems are found, qualification of the affected data points will be recommended. Upon determination of the level of quality for each data point, interpretation of the data can then be performed. The following sections describe the data quality evaluation and data qualification processes.

1.9.3.1 <u>Data Quality Evaluation</u> - The objective of the evaluation the quality of the chemical data is to determine qualifications of the data are necessary. This evaluation will be based upon the evaluation of the laboratory QC data, the field QC data, and the project DQOs presented in Section 1.4. The first step will be to perform an evaluation of the laboratory QC data, a process often termed "data validation." This will provide a rating of the quality of each data point produced by the laboratory. second part will provide an overall rating of each data point based upon the field QC data. The final step in the evaluation will compare the quality of the data acquired to the project's DQOs to determine whether the data are useful. These three steps are described in the following sub-sections. Each step will be completely documented. The overall goal of the data quality evaluation is to determine whether the data can be used to satisfy the objectives of the project.

Evaluation of Laboratory QC Data - Laboratory QC data for an analytical parameter fall within one of eight categories for evaluation purposes. These categories are listed in Table 1-12 in the order in which they will be considered. All QC data provided will be evaluated against the criteria established by each method after modification as presented in this document to achieve the objectives of this project. Each review will be completely documented to indicate the criteria used and the results and recommendations of the evaluation. For this investigation the evaluation of the calibration will be performed by the Law Environmental National Laboratory in Pensacola. This laboratory verification of method calibration is in compliance with an AFCEE Level I data package.

Evaluation of Field OC Data - Field QC data for an analytical parameter fall within one of eight categories for evaluation purposes. These categories are also listed in Table 1-12 in the order in which they will be considered. QC data provided will be evaluated to assure the objectives of this project are achieved. Each review will be completely documented to indicate the criteria used and the results and recommendations of the evaluation.

<u>Usability Determination</u> - Once the laboratory and field QC data have been evaluated, the uncertainty associated with each data point can be estimated. The estimated accuracy and precision of each data point can then be compared to the data quality objectives of the project to determine its usefulness for evaluating the site. Recommendations for the qualification of a data point can also be made when necessary.

Precision will be determined by evaluation of the RPDs for the laboratory and field duplicates. The laboratory and field blank data, MS/MSD and surrogate (if applicable) recoveries, and other applicable QC data will be used to determine the accuracy of the data. The evaluator will use their judgement based upon

TABLE 1-12

DATA EVALUATION CATEGORIES RCRA Facility Investigation Carswell Air Force Base, Texas

187 81

	CATEGORY	QC DATA EVALUATED
LAE	SORATORY QC:	
1.	Sample Integrity	Sample container condition; preservation performance and applicability; condition upon receipt at laboratory
2.	Instrument Set-up/Method Applicability	Correct method; detection/quantitation limits achieved; method applicability for analytes and matrix, instrument set-up
3.	Calibration Accuracy and Precision *	Standard preparation; accuracy verification, precision achieved
4.	Calibration Stability *	Stability verification; frequency of performance
5.	Laboratory Contamination	Laboratory blanks
6.	Method Accuracy and Precision	Method accuracy and precision data
7.	Sample Preparation	Procedures; holding times
8.	Sample Analysis	Procedures and sequences; holding times; sample- specific accuracy and precision; interferences; dilutions
FIE	<u>-D QC</u> :	
1.	Sample Integrity	Sample container condition; preservation performance and applicability; condition prior to shipment; custody
2.	Location Installation	Sampling location installation
3.	Sampling Procedures	Sampling protocols
4.	Contamination From Sampling	Rinsates
5.	Contamination From Site	Field blanks (volatile organics only)
6.	Contamination From Shipping	Trip blanks (volatile organics only)
7.	Sampling Precision	Field duplicates
8.	Miscellaneous +	Blind QC samples; performance evaluation data; split sample data

^{*} These categories are assumed to be correct and will not be provided for evaluation.

⁺ No blind data for QC samples, blind performance evaluation samples, or split samples will be collected.

established principles and the guidelines described in this document in the assignment of qualifications to the data.

parameter will The completeness be evaluated after t.he determination of the usability of each data point; it will be expressed in quantitative terms and then compared to the project objectives to determine whether enough data were collected. Representativeness will be determined through an evaluation of sampling procedures and locations utilized and will be expressed in Comparability will be determined by the qualitative terms. evaluation of data quality analytical methodologies, reporting units, and the traceability of standards and will also be expressed in qualitative terms.

1.9.3.2 <u>Data Qualification</u> - Each data point will essentially be graded as falling into one of the following categories:

- Usable as reported
- Usable with qualifications
- · Unusable

These categories correspond to the three established for the DQOs in Section 1.4. Data for which the laboratory and field QC data are all within acceptance limits will be assigned the grade "usable as reported." Data for which slight QC problems are indicated but the QC data are still within the action limits will be assigned the grade "usable with qualifications." These data may still be used if the QC problems are not excessive. Data for which the corresponding QC data are outside the action limits will be assigned the grade "unusable" and will not be used. Each data point will receive a flag indicating its level of quality. The flags to be used are presented in Table 1-13.

DATA QUALIFICATION FLAGS RCRA Facility Investigation Carswell AFB, Texas

FLAG	POSITIVE RESULTS	NEGATIVE RESULTS
FLAGS FOR DA	ATA WITHIN ACCEPTANCE LIMITS (Usable a	s Reported)
(no flag)	{Use datum without qualification}	{Use datum without qualification}
FLAGS FOR DA	ATA WITHIN ACTION LIMITS (Usable With Q	ualifications)
J	Estimated quantitation based upon QC data	Estimated quantitation based upon QC data
JB	Estimated quantitation: possible biased high or false positive based upon blank data	(Not applicable)
JH	Estimated quantitation-possibly biased high based upon QC data	(Not applicable)
JL	Estimated quantitation-possibly biased low based upon QC data	Possible false negative based upon QC data
Jd	Estimated result due to dilution	Reporting limit raised due to dilution
FLAGS FOR D	ATA OUTSIDE OF ACTION LIMITS (Unusable)	
R	Datum rejected based upon QC data: do not use	Datum rejected based upon QC data: do not use
MISCELLANEC	DUS FLAGS	
t	Tentatively identified compound; identity not confirmed with standard and quantitation estimated*	(Not applicable)

^{*} Applicable to GC/MS data only

1.9.4 Data Reporting

Data reports will be included in the technical reports preparation. The data will be presented as tables or in the appendices of the report. Tables will include the following information:

- · Sampling dates
- · Extraction and analysis dates
- Surrogate recovery (if applicable)
- MS/MSD results
- · Duplicate/replicate results
- · Rinsate results
- · Positive results
- · Field characterization data (pH, SC, temp)
- · Control limits (surrogates, MS/MSD, duplicates)

1.10 INTERNAL QUALITY CONTROL CHECKS

The following sections describe the quality control checks employed in the field and laboratory.

1.10.1 Field Quality Control

Quality control of field measurements will be utilized through the calibration of instruments. The control parameters, control limits, and corrective actions are outlined in Section 2.4.

1.10.2 <u>Laboratory Quality Control</u>

The minimum requirements of the laboratory quality control consist of an initial demonstration of laboratory capability and an ongoing analysis of quality control samples to evaluate and document data quality. The laboratory must maintain records to document the

quality of the data generated. Ongoing data quality checks are compared with established performance criteria to determine if the results of analyses meet the performance characteristics of the method. When results of quality control samples indicate atypical method performance, a quality control check standard must be analyzed to confirm that the measurements were performed in an incontrol mode of operation. Calibration stability will be assessed and documented utilizing the procedures and at the frequency specified in each method.

Before processing any samples, the analyst should demonstrate, through the analysis of a reagent water blank, that interferences from the analytical system, glassware, and reagents are under control. Each time a set of samples is extracted or there is a change in reagents, whichever is more frequent, a reagent water blank should be processed as a safeguard against chronic laboratory contamination. The blank samples should be carried through the stages of the sample preparation and measurement steps.

If any analyte fails the acceptance criteria for recovery, a QC check standard for each analyte that failed must be prepared and analyzed. As part of the QC program for the laboratory, method accuracy for each matrix studied must be assessed and records must be maintained.

The types and numbers of laboratory quality control samples to be used are presented in Tables 1-14 through 1-18 by matrix and parameters. They are defined as follows:

Method blanks. Method blanks consist of organic-free or deionized water that is carried through the analytical scheme like a sample. They serve to measure contamination associated with laboratory storage, preparation, or instrumentation. For most analyses, a method blank is analyzed for each batch and at a frequency of 1 per 20 samples if more than 20 samples are run in a given batch. If

TABLE 1-14

SAMPLING AND ANALYSIS PLAN SUMMARY: SUB-SURFACE SOIL FROM BORINGS RCRA Facility Investigation - Carswell Air Force Base, Texas

	TOTAL	IAB SAMPLES	57	9	. 25 0		. o .	٥	<u>.</u> 6.	ا م	ن ۱	7
	2ND (c)	CONFIRM. ANALYSES	- •	- 2		-	N 00		· • •		o .	0
		Spiked Blank (b)	- 0	· -	- 0	-	0	-	0	-	· o	0
	LAB QC SAMPLES	Lab Duplicate	0 0	0	00	0	00	0	00	0	0	0
	LAB	MSD	- 0	-	- 0	-	- 0	-	-0	-	-	-
),	W	- 0	-	- 0	-	-0	-	- 0	-	-	-
ATOT	NO E	SAMPLES	ထေ	16	∞ ∞	16	യയ	12	യയ	12	വ	ر م
	SAMPLES	LOCATION	W 4		0.4		Ο 4		o 4		9	
TOTAL	NO.	LOCATIONS	o –	8	8 +	8	c +	3	0 -	ဇ	ო	ဇ
		PARAMETER	Volatile Organics (SW 8240) Unnamed Stream: Downgradient (d) Unnamed Stream: Upgradient (e)	TOTALS	Normatic Volatiles (SW 8020) Unnamed Stream: Downgradient Unnamed Stream: Upgradient	TOTALS	<u>Total Recoverable</u> Petroleum Hydrocarbons (E 418.1) Unnamed Stream: Downgradient Unnamed Stream: Upgradient	TOTALS	ICP Screen for Metak (gas 3050/6010) Unnamed Stream: Downgradient Unnamed Stream: Upgradient	TOTALS	Toxicity Characteristic Leaching Procedure (SW 1311) (f) Unnamed Stream	TOTALS

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan.

(a) Volatile Organics only; number to be initiated will depend upon number of shipments.

(b) Estimated; number to be determined by batch preparation.

(c) Methods SW 8240 and SW 8020 will have 2nd column confirmation performed on all samples exhibiting positive results.

(d) Soil samples from oil/water separator and Unnamed Stream monitoring well locations.

(e) Soil samples from single upgradient monitoring well installation.

(f) TCLP analyses will be performed on the containerized waste material generated from the drill cuttings.

TABLE 1-15

ANALYSIS SUMMARY: GROUND WATER - MONITORING WELLS RCRA Facility Investigation - Carswell Air Force Base, Texas

		TOTAL	TOTAL NO.	TOTAL NO.		LAB QC	LAB QC SAMPLES	1 1	2ND (b) COLUMN	TOTAL NO.
PARAMETER		FIELD	SAMPLING	FIELD	WS	MSD	Lab Duplicate	Spiked Blank (a)	CONFIRM. ANALYSES	LAB
Volatile Organics (SW 8240) Unnamed Stream		თ	ო	27	ო	ო	0	ო	0	36
	TOTALS	6	ဇ	27	က	ဇ	0	က	0	36
Total Recoverable Petroleum Hydrocarbons (418.1) Unnamed Stream		თ	ဇ	27	ო	ო	0	ო	0	36
	TOTALS	o	က	27	က	3	0	က	0	36
Oil and Grease (SW 9071) Unnamed Stream		თ	ო	27	ო	ო	0	ო	0	36
	TOTALS	o	က	27	ဧ	က	0	က	0	36
Screen for 24 Metals (SW 6010) Unnamed Stream: Total Unnamed Stream: Dissolved		თთ	ოო	27 27	ოო	ო ო	0 0	ო ო	0 0	36 36
	TOTALS	18	9	54	9	9	0	9	0	72
Arsenic (SW 7060) Unnamed Stream: Total Unnamed Stream: Dissolved		თთ	ოო	27 27	ღ ღ	ო ო	0 0	ოო	0 0	36 36
	TOTALS	8	9	54	မ	9	0	9	0	72

38

187

TABLE 1-15

ANALYSIS SUMMARY: GROUND WATER - MONITORING WELLS RCRA Facility Investigation - Carswell Air Force Base, Texas

		TOTAL NO.	TOTAL NO.	TOTAL NO.		LAB QC	LAB QC SAMPLES		2ND (b) COLUMN	TOTAL NO.
PARAMETER		FIELD	SAMPLING	FIELD SAMPLES	MS	MSD	Lab Duplicate	Spiked Blank (a)	CONFIRM. ANALYSES	LAB
Mercury (SW 7470) Unnamed Stream: Total Unnamed Stream: Dissolved		თთ	ღღ	27 27	ღღ	ი ი	00	ოო	00	36 36
	TOTALS	18	9	54	9	9	0	9	0	72
Selenium (SW 7740) Unnamed Stream: Total Unnamed Stream: Dissolved		തത	ო ო	27 27	ღღ	ო ო	00	ო ო	00	36 36
	TOTALS	18	ဖ	54	9	9	0	9	0	72
Lead (SW 7421) Unnamed Stream: Total Unnamed Stream: Dissolved		തത	ო ო	27 27	ღღ	ო ო	00	ო ო	00	36 36
	TOTALS	18	9	54	ဖ	ဖ	0	9	0	72
Toxicity Characteristic Leaching Proœdure (SW 1311) Unnamed Stream (c)		5	-	52	-	-	0	0	0	7
	TOTALS	S	-	Ŋ	-	· -	0	0	0	7

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan.

⁽a) Estimated; number to be determined by preparation batch.(b) Method SW 8240 will have 2nd column confirmation performed on all samples exhibiting positive results.(c) The purge water will be analyzed for the parameters of concern in the TCLP list.

TABLE 1-16

SAMPLING AND ANALYSIS PLAN SUMMARY: GROUND WATER - FIELD SCREEN RCRA Facility Investigation - Carswell Air Force Base, Texas

		TOTAL NO.	TOTAL NO.		LAB QC	AB QC SAMPLES		2ND (c)	TOTAL
		FIELD	FIELD	1778		Lab	Spiked	CONFIRM	YB :
PARAMETER		LOCATIONS	SAMPLES	MS	MSD	Duplicate	Blank (b)	ANALYSES	SAMPLES
Aromatic Volatiles (SW 8020)									
POL Tank Farm		22	26	8	2	0	-	10	4
Reserve (d)		က	22	-	-	0	0	-	80
	TOTALS	25	31	တ	က	0	-	-	49
Petroleum Hydrocarbons (SW 8015M)									
POL Tank Farm		22	26	8	8	0	-	9	51
Reserve (d)		ဇာ	23	•	-	0	-	-	٣
	TOTALS	25	31	ဇ	ဇ	0	2	-	29
Lead (SW 6010)									
POL Tank Farm		22	24	2	2	0	-	0	29
Reserve (d)		ဇာ	က	1	-	0	0	0	Ŋ
	TOTALS	25	27	၈	ဧ	0	-	0	34

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan.

(a) Volatile Organics only; number to be initiated will depend upon number of shipments.

(b) Methods SW 8020 and SW 8015M will have ten percent of all samples sent to an off-site laboratory for confirmatory analyses.(c) Three ground-water sampling locations will be reserved for discretionary use during the implementation of the field work.

ANALYSIS SUMMARY: SURFACE WATER

RCRA Facility Investigation - Carswell Air Force Base, Texas

TOTAL NO.	LAB SAMPLES	ဖ	မ	ဖ	9	ဖ	g
2ND (b) COLUMN	CONFIRM. ANALYSES	0	0	0	0	0	0
	Spiked Blank (a)	-	-	-	-	-	-
LAB QC SAMPLES	Lab Duplicate	0	0	0	0	0	0
LAB QC	MSD	-	-	-		-	-
	WS WS	-	-	-	-		-
TOTAL NO.	FIELD	ო	က	ო	က	ო	က
TOTAL NO.	FIELD	ო	က	ო	ဇာ	ო	ဇာ
			TOTALS		TOTALS		TOTALS
	PARAMETER	Volatile Organics (SW 8240) Unnamed Stream		Total Recoverable Petroleum Hydrocarbons (418.1) Unnamed Stream		Oil and Grease (SW 9071) Unnamed Stream	

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan.

187 90

⁽a) Estimated; number to be determined by preparation batch.(b) Method SW 8240 will have 2nd column confirmation performed on all samples exhibiting positive results.

TABLE 1-18

RCRA Facility Investigation - Carswell Air Force Base, Texas **ANALYSIS SUMMARY: SEDIMENT**

TOTAL NO.	LAB SAMPLES	တ
	Spiked Blank (a)	- -
LAB QC SAMPLES	Lab Duplicate	0 0
LAB Q	MSD	- -
	MS	
TOTAL NO.	FIELD	ო ო
TOTAL NO.	HELD LOCATIONS	ო ო
	PARAMETER	ICP Screen for Metals (SW 3050/6010) Unnamed Stream TOTALS

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan. (a) Estimated; number to be determined by preparation batch.

the analyte of interest is above the reporting detection limit, corrective action should be taken except for common solvents such as methylene chloride, acetone, toluene, 2-butanone and phthalates.

<u>Sample blanks</u>. Sample blanks are used when characteristics like color or turbidity interfere with a determination. In a spectrophotometric method, for example, the natural absorbance of the sample is measured and subtracted from the absorbance of the developed sample. Sample blanks are run only as necessary.

<u>Calibration blanks</u>. Calibration blanks are prepared with standards to create a calibration curve. They differ from the other standards only by the absence of analyte and provide the "zero-point" for the curve.

<u>Internal standards</u>. Internal standards are measured amounts of certain compounds added after preparation or extraction of a sample. They are used in an internal standard calibration method to correct sample results suffering from capillary column injection losses, purging losses, or viscosity effects. Internal standard calibration is currently used for volatile organics, chlorinated pesticides and GC/MS extractables.

<u>Surrogates</u>. Surrogates are measured amounts of certain compounds added before preparation or extraction of a sample. The recovery of a surrogate is measured to determine systematic extraction problems. Surrogates are added to all samples analyzed for chlorinated pesticides, GC/MS extractables and volatiles, and GC volatiles.

<u>Spikes</u>. Spikes are aliquots of samples to which known amounts of analyte have been added. They are subjected to the sample preparation or extraction procedure and analyzed as samples. The stock solutions used for spiking are purchased or prepared independently of calibration standards.

The spike recovery measures the effects of interferences in the sample matrix, and reflects the accuracy of the determination. Spike recoveries are calculated as follows:

Check Standard Method Standard Check Sample

Percent Recovery = $\frac{\text{Observed}}{\text{Expected}} \times 100$

Matrix Spike

Percent Recovery = $\frac{SSR - SR}{SA}$ x 100

where:

SSR = spike sample result
SR = sample result
SA = spike added from spiking mix

Spikes are prepared and analyzed for all method analytes with each batch and at a frequency of at least 1 MS and 1 check standard per 20 samples if more than 20 samples are run in a given batch, or as required by the specific analytical method.

Spike recoveries are stored in the laboratory database and are retrievable for statistical analysis. Laboratory control limits are calculated for individual matrix types when 20 data points become available.

<u>Duplicate</u> or <u>Duplicate</u> Spikes. Duplicate spikes are additional spiked aliquots of samples subjected to the same preparation and analytical scheme as the original spike sample. The relative percent difference (RPD) between duplicates or duplicate spikes measures the precision of a given analysis. RPDs are calculated as follows:

$$RPD = \frac{R1 - R2}{Rav} \times 100$$

%RPD =
$$\frac{S1 - S2}{Sav} \times 100$$

where:

R1 and R2 = duplicate determinations of the analyte in the sample

S1 and S2 = the observed concentrations of analyte in the spike and its duplicate

Rav = the average determination of the analyte concentration in the original sample

Sav = the average of the observed analyte concentrations in the spike and its duplicate

Duplicates or duplicate spikes are prepared and analyzed for each method analyte with each batch, or at a frequency of at least 1 per 20 samples if more than 20 samples are run in a given batch.

RPDs are stored in the laboratory database and are retrievable for statistical analysis.

Laboratory Control Standards. Laboratory Control Standards (LCSs) and Quality Control Check Samples (QCCSs) are aliquots of organic-free or deionized water to which known amounts of analyte have been added. They are subjected to the sample preparation or extraction procedure and analyzed as samples. The stock solutions used for LCSs are purchased or prepared independently of calibration standards. The LCS recovery tests the function of analytical methods and equipment. For inorganic and metals analyses, the percent recovery for LCSs is compared to method specific criteria, and the analytical system is considered to be in control when these analyte specific criteria are met. When a result of an organic method aqueous matrix spike and/or matrix spike duplicate indicates atypical method performance, a quality control check standard will

be analyzed to confirm that the measurements were performed in an in-control mode of operation for that analyte.

The acceptance criteria for the LCS is a recovery range of 80 to 120 percent. The acceptance criteria for the QCCS is stated in each method.

LCSs are prepared and analyzed for all method analytes with each batch or at a frequency of 1 per 20 samples if more than 20 samples are run in a given batch. Laboratory control limits are calculated when 20 data points become available.

The LCS is used to monitor overall performance of all steps in analysis, including sample preparation. If the LCS results fall within 20 percent of the expected or true value calibration curve, this verifies that instrument performance, calibration and sample preparation are satisfactory. When the LCS is used in conjunction with matrix spikes, matrix spike recovery evaluation can be better interpreted as either matrix interference, sample preparation error or matrix spiking solution preparation error.

1.10.3 Control Limits

Control limits for this project are method specific and laboratory established. Control limits for each analytical method will be experimentally established and reevaluated at regular intervals. Control limits are developed by the laboratory based on historical data. If historical data are not complete then control limits are set based on those established by the method. Appendix A presents the control limits for each analytical method including frequency, acceptance criteria, and corrective action.

1.11 PERFORMANCE AND SYSTEM AUDITS

Law views quality assurance as the means by which the effectiveness and quality of its various participating departments are gauged as they carry on day-to-day operations under the QA/QC program. The major goals associated with the QA/QC program are listed below:

- QA reviews should help ensure compliance with mandated QC procedures.
- QA reviews provide a structured means of communicating problems between the technical and administrative portions of the company.
- QA procedures are designed to ensure operating regularity between the various branches of Law.
- QA audits provide a mechanism by which our QC procedures are constantly being reviewed and updated in an orderly fashion.

1.11.1 Systems Audits

Systems audits are qualitative evaluations of each component of field and laboratory QC measurement systems. A systems audit will be performed periodically and will consist of inspecting the following procedures:

- · Sampling
- · Sample custody
- · Sample storage and preservation
- · Standard preparation
- · Sample preparation
- Analytical methodology

- · Data management
- · Preventative maintenance
- · Personnel qualifications
- Corrective actions, reporting and documentation of out-ofcontrol events

187 97

· Recordkeeping

Law will submit to on-site external systems audits by AFCEE or any other appropriate regulating agency.

1.11.2 Performance Audits

A performance audit is a quantitative evaluation of a measurement system. Law participates in the following performance evaluation programs:

- · EPA, Environmental Monitoring Support Laboratory Cincinnati
- · Florida Department of Environmental Regulation (DER)
- · EPA Inorganic and Organic CLP
- · U.S. Army Corps of Engineers, Missouri River Division
- Florida Department of Health and Rehabilitative Services (FDHRS)
- U.S. Air Force Performance Audit conducted by MITRE, in support of AFCEE

Periodically, blind QC samples are entered into the samples flow scheme. Since blind QC samples are difficult to enter into the laboratory without the knowledge of the analysts, the QA Manager relies heavily on the performance evaluation programs by outside sources and QC samples such as matrix spikes, duplicates and surrogates.

Law Environmental National Laboratories located in Pensacola, Florida currently holds certifications with several states for various matrices and parameters. They are also certified with the COE, USAF, and USEPA.

1.12 PREVENTIVE MAINTENANCE

Equipment maintenance is the responsibility of the analyst and the Department Manager. All repairs and/or modifications are recorded in bound maintenance logbooks. Daily equipment checks include visual and/or manual inspections of cooling fans, pumps, indicator readings, detectors, gas supplies, and other method-specific inspections. Service schedules are established for performing routine preventive maintenance on all major equipment. Tables 1-19 through 1-23 list the maintenance required for each instrument.

1.13 <u>FIELD AND LABORATORY PROCEDURES USED TO ASSESS DATA QUALITY</u> INDICATORS

1.13.1 Formulas

Accuracy - Accuracy is a measure of the bias in a system. Accuracy is defined as the degree of agreement of a measurement with an accepted reference or true value. To determine the accuracy of an analytical method, a sample spiking program will be conducted. The results of sample spiking will be used to calculate the quality control parameter for accuracy evaluation, the Percent Recovery (%R). The Percent Recovery is defined as the observed concentration, minus the sample concentration, divided by the true concentration of the spike, multiplied by 100.

TABLE 1-19

ROUTINE MAINTENANCE SCHEDULE FOR GC Carswell Air Force Base, Texas

Maintenance	Frequency
Check carrier gas supply	Daily
Check head pressure	Daily
Change septums	As needed
Change carrier gas filters	As needed
Check baseline and detector response at highest sensitivity	Quarterly
Air dust electronics and main frame	Annually

ROUTINE MAINTENANCE SCHEDULE FOR GC/MS Carswell Air Force Base, Texas

	Maintenance	Frequency			
1.	System Diagnostics				
	a. Scan box I/O test	Daily			
	b. Scan box DMA test	Daily			
	c. MS I/O test	Daily			
	d. Filament test	Daily			
	e. RF power test	Daily			
	f. Electromultiplier test	Daily			
	g. Background signal noise test	Daily			
2.	Vacuum/Carrier Flow Check				
	a. Column headpressure check	Daily			
	b. Vacuum check	Daily			
	c. Replace septum	As needed			
	d. Leak check and tighten fittings	As needed			
3.	Instrument Tune Check				
	a. BFB/DFTPP tune check	Daily			
	b. PFTBA tuning	As needed			
4.	Calibration Check				
	a. VOA standards check	Daily			
	b. BNA standards check	Daily			
	Replace inlet liner	As needed			
5.	Clean source	As needed			
6.	Clean quadrupole rods	As needed			
7.	Replace column	As needed			

ROUTINE MAINTENANCE SCHEDULE FOR ICP Carswell Air Force Base, Texas

	MAINTENANCE	FREQUENCY
Check Argon pressu	ire at tank and at nebulizer.	Daily
2. Inspect cooling wat	er supply.	Daily
3. Inspect vacuum pur	np.	Daily
a. Oil level b. Vacuum monoo	chromator gauge	
4. Inspect peristaltic p	ump windings and capillary tubing.	Daily
5. Check ICP ignition s	sequence.	Daily
a. Argon to torchb. Preignition discc. Ignition	harge	
 Disassemble and cloassembly. 	ean ICP nebulizer, spray chamber and torch	Weekly
Check peak resoluti profile routines and	on and monochromator stability using internal Hg lamp.	Weekly
8. Change pump wind	ings and all capillary tubing to nebulizer.	Monthly
9. Change vacuum pu	mp oil.	Monthly

ROUTINE MAINTENANCE SCHEDULE FOR AA SPECTROPHOTOMETER Carswell Air Force Base, Texas

Maintenance

- 1. Contact cylinders, tube, and platform should be cleaned and checked each day before running samples. Use cotton swab and methanol for cleaning.
- 2. Contact cylinders should be checked for cracks and pitting. Tubes should be checked for pitting, peeling pryolitic coating and burn marks around sample port hole. Any of the above indicate a worn tube or cylinder.
- 3. Furnace/spectrometer windows should be checked and cleaned daily.
- 4. Check coolant level in recirculator and temperature setting daily.
- 5. Spectroscopy lab should be wet mopped, counters dusted, and exterior of instruments cleaned on a weekly basis to ensure a dust-free environment.
- 6. Consult instrument operations manual for further maintenance instructions.
- 7. All maintenance is to be recorded in the Maintenance Log Book.
- 8. Argon gas pressure to furnace should be 60 psi. Check regulator to ensure proper pressure.

TABLE 1-23

ROUTINE MAINTENANCE SCHEDULE FOR IR SPECTROPHOTOMETER Carswell Air Force Base, Texas

187103

MAINTENANCE	FREQUENCY	
1. Check air filter.	Depends on use	
2. Remove cuvette from sample compartment.	Daily	
3. Keep instrument clean from spills.	Daily	

$$R = \frac{X - T}{K} \times 100$$

187104

where:

T = analytical result from the unspiked aliquot

K = known value of the spike

%R = percent recovery

To determine accuracy, surrogate, matrix spike and matrix spike duplicates (MS/MSD), and internal standards will be analyzed. The control limits will be based on the mean percent recovery plus or minus 3 standard deviations of the mean using a population of 20 recovery values.

<u>Precision</u> - Precision is the measure of mutual agreement among individual measurements of the same property, usually under prescribed similar conditions. Precision is best expressed in terms of standard deviation or Relative Percent Difference (RPD) Precision is inferred through the use of duplicate samples. RPD for each component is calculated using the following equation:

$$RPD = \frac{A - B}{(A + B)/2} \times 100$$

where:

A = replicate value 1

B = replicate value 2

RPD = relative percent difference

The calculated Percent Recovery and RPD will be summarized. The RPD data will be used to evaluate the long term precision of the analytical method.

To determine precision, matrix spikes and matrix spike duplicates will be analyzed. The control limits will be based on a population of ten RPD values. They are calculated by determining the mean RPD plus three times the standard deviation for the upper limit and zero as the lower limit.

<u>Completeness</u> - Completeness is a measure of the amount of valid data obtained from a measurement system compared to the amount that was expected to be obtained under correct or normal conditions. The result is expressed as a percentage.

Percent completeness = $\frac{\text{Number of valid measurements}}{\text{Total number of measurements}} \times 100$

The percent completeness for this project is 90 percent.

1.13.2 Control Limits

Control limits for analytical methods are presented in Appendix A. Control limits are developed by the laboratory based on historical data. If historical data are not complete then control limits are set based on those established by the method.

1.13.3 Documentation

Completeness and comparability of data are insured by adherence to a standard data-set protocol and checklist of data required to be available on laboratory report sheets for each type of analysis conducted. Each parameter data book must contain all data and calculations associated with each independent determination. These include such things as sample weight(s), dilution factor(s), applicable determinative measurements such as titration values,

spectrometer readings, injection quantities, and standard(s) identity and concentrations, as well as all calculations related to each final value reported. Each laboratory report sheet will be checked and initialed by a second competent scientist other than the person who did the analysis to insure completeness of data and correctness of all calculations in the report. For the occasional project involving very critical samples on which serious action is contemplated, the entire set of analyses on the same sample(s) will be independently audited by a special project quality assurance officer, (for example, the analysis of drinking water samples from residential wells where compounds were found to exceed the MCL). Samples collected at Carswell AFB are not considered to be in this Work sheets, chromatograms, spectra, etc., associated with every analysis will be present in the parameter data book. These will give instrument operation parameters and details of instrument set-up, such as columns used, mobile and stationary phases and concentrations, temperatures, detectors, spectrometer type, wavelengths, etc., as applicable for the type analysis. Work sheets will bear a unique laboratory number identifying that sample, relating the laboratory data sheets to the sample. sheet will be recorded with dates and times and bear the laboratory analyst's initials.

1.14 CORRECTIVE ACTION

An effective QC program requires rapid and thorough correction of the QC problems. Rapid corrective action (CA) minimizes the possibility of questionable data on any project. The need for corrective action originates when an inadequacy is found in the method of analysis (e.g., inappropriate calibration) or a determinate error occurs (e.g., calibration error due to standards failure). Failures of the first kind are precluded by LENL and Regulator/Contractee audits which evaluate analytical SOPs. The analytical SOPs incorporate mechanisms to detect the existence of determinate errors and specify the procedures to correct them.

Depending on the nature of the CA, it is classified as one of two types, immediate and long-term. Immediate corrective actions are the correction of procedures or repair of instrumentation that is working improperly. Long-term corrective actions are the correction of systematic errors, such as the detector on a GC becoming dirty and losing sensitivity. Corrective actions associated with field activities are presented in Section 2.4.3.

1.14.1 Response

Many times the source of a nonsystematic problem is obvious to the analyst and can be corrected immediately. Immediate corrective action routinely made by field technicians or laboratory analysts should be documented as normal operating procedures in instrument log books or personal notebooks. The Supervisor and analyst should compile a list of commonly encountered problems and the appropriate routine corrective actions (in addition to manufacturer's troubleshooting guides). The Operations Manager and QA/QC Coordinator are responsible for approving all corrective actions. Appendix A lists the corrective actions necessary for each analytical method.

1.14.2 Reestablishment of Control

Corrective action is not complete until the problem has been effectively and permanently solved. Follow-up action to ensure that the problem remains corrected is a vitally important step in the corrective action procedure. Routine corrective actions, such as recalibrating the instrument, are incorporated into the Standard Operating Procedures (SOPs). Major corrective actions, such as a systems failure, are handled in the following manner: Once a problem has been technically defined, the Operations Manager and the QA/QC Coordinator discuss the problem and jointly take the following steps:

- 1. Determine that specific corrective action is needed to eliminate the problem and assign responsibility for investigating, implementing, and documenting the situation.
- 2. Set a time schedule for determining the required action.
- 3. Assign responsibility and time schedule to implement the desired action.
- 4. Establish desired effectiveness of the corrective action and implement the correction.
- 5. Verify that the corrective action has eliminated the problem and document.

1.14.3 Documentation

To provide a complete record of QC activities, QC problems and corrective actions applied must be documented. Historical records assist laboratory management in identifying long-term corrective actions. such as personnel training, replacement of instrumentation, improvement of sampling procedures, Corrective action documentation forms for laboratory operations are Appendix B. Corrective action documentation in associated with field activities are recorded on Daily Quality Control Forms. These forms are presented in Section 2.5.

1.15 **QUALITY ASSURANCE REPORTS**

The management is informed of QA activities in three ways: (1) by immediate verbal notification of QA problems, (2) by interim QA

reports, and (3) written QA reports. The following subsections discuss the reporting procedures to be followed and report contents.

1.15.1 Reporting Procedure

An Analytical Data Informal Technical Information Report (ITIR) will be prepared by the Project Chemist, reviewed by the Project Manager and submitted by Law Environmental Government Services Division at the end of the sampling and analysis activities to the AFCEE TC. The data will also be submitted to the USAF TPM in a format compatible with the USAF's Installation Restoration Program Information Management System (IRPIMS) data base.

1.15.2 Report Content

The format of the Analytical Data ITIR will substantially follow the format specified by the USAF (AFCEE, 1991). The format of the IRPIMS deliverable will correspond exactly to the format specified by the USAF (AFCEE, 1991; AFCEE, 1993a; AFCEE, 1993b). The internal management of these deliverables is performed by a QA audit between the LENL-P Chemical Lab Manager and the LEGS Project Manager (see Figure 1-5). The QA audit reporting function is facilitated by direct correspondence between the LEGS Chemistry Program Lab Coordinator and the lab.

2.0 FIELD SAMPLING PLAN (FSP)

Field tasks to be conducted at the Carswell AFB RFI sites are explained in detail in the following text.

2.1 FIELD OPERATIONS

The field investigations at the Unnamed Stream and POL Tank Farm sites which are located in the east area of Carswell AFB (Figures 2-1 and 2-2) will include the following activities:

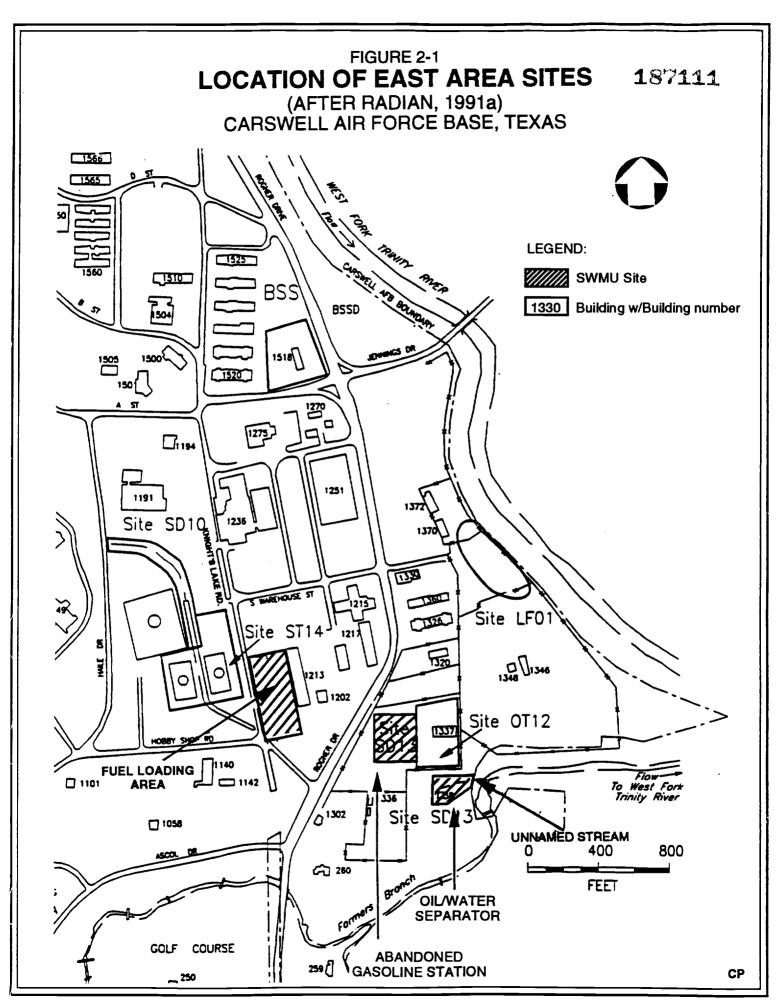
- · Geophysical surveys
- · Geochemical survey
- · Installation of new monitoring wells
- · Aquifer testing
- Soil, sediment, ground-water sampling (three rounds), and surface water sampling

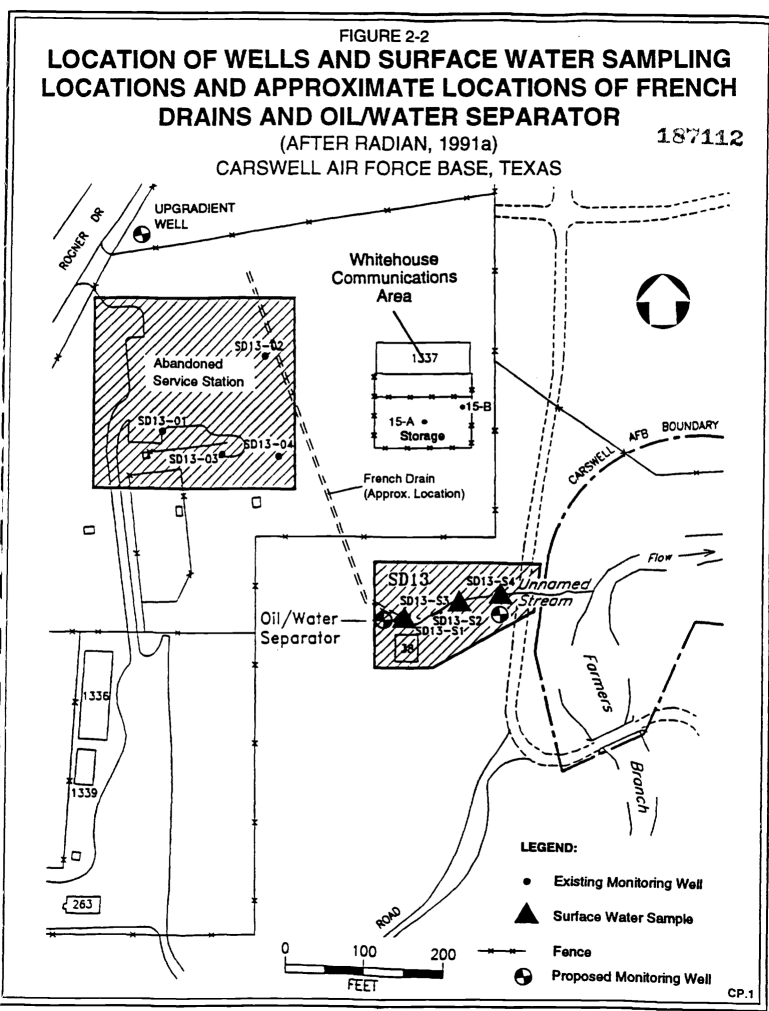
The proposed geochemical locations for the POL Tank Farm are presented in Figure 2-3. The proposed monitoring well location for the Unnamed Stream is presented in Figure 2-2. The methods for conducting the field operations and activities are discussed in the following sections.

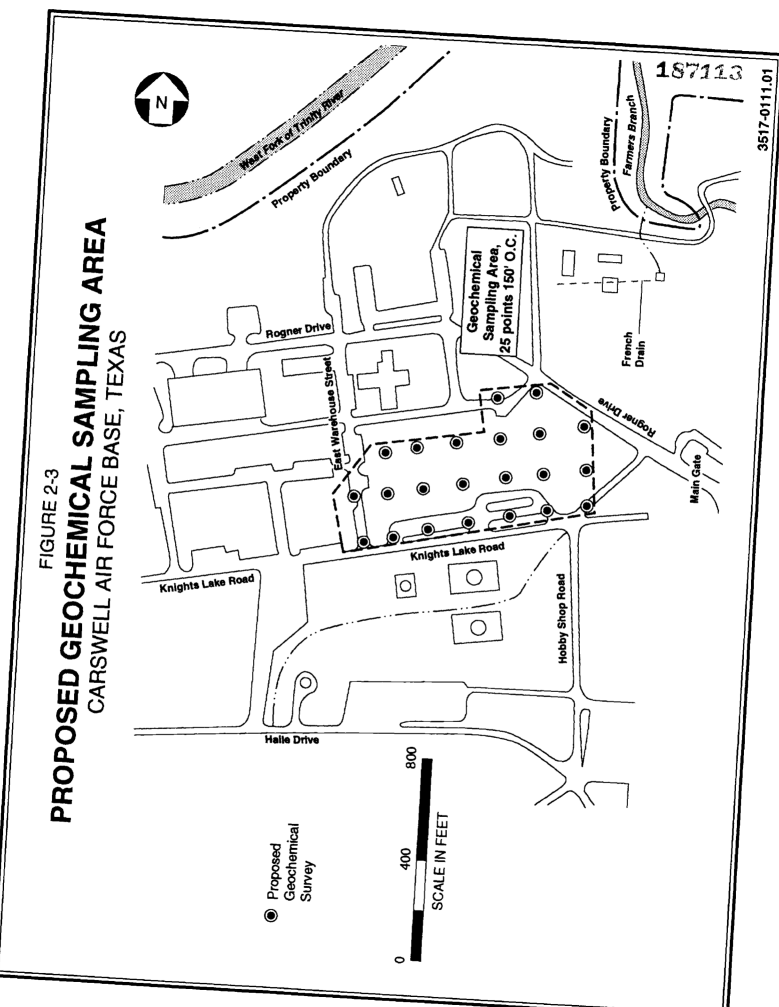
2.1.1 Site Reconnaissance, Preparation and Restoration

The two sites under this RFI were visited on April 14th, 1993, by representatives of Law, AFCEE, and Carswell AFB. During the site visit, AFCEE representatives outlined the RFI field tasks at each site.

Prior to commencement of field operations, all sites will be inspected and new monitoring well locations will be staked. The sites will also be inspected for access routes.







Site preparation will include removal of obstructions (if needed) and utility clearances by base personnel. Specific information concerning field office space, decontamination area, and drum staging areas will be coordinated with base personnel.

Upon completion of field activities, each site will be restored to a condition that as closely as possible approximates the condition of the site prior to field activities.

2.1.2 <u>Surface Geophysical Survey</u>

A geophysical survey will be performed at both sites to help characterize the near surface conditions and to locate utilities and abandoned tanks and pipechases. A single type of geophysical survey (ground penetrating radar) will be performed at Carswell AFB during the field work.

2.1.2.1 Ground Penetrating Radar - GPR is a geophysical technique which can provide high resolution data on surficial geology. The technique is used in general to map shallow bedrock, soil and water table features, and locate underground pipes and tanks. At hazardous waste sites, ground penetrating radar is also used to locate trenches, lagoons, buried drums, contaminant plumes and other structural and contaminant-related features. The depth of penetration is dependent upon the types of soil and the electrical properties of the subsurface. In silts and clays the depth of penetration may be on the order of only a few feet, while in dry sands the depth of penetration may extend to tens of feet.

A high frequency signal (typically in the 100 to 1000 Mhz range) is transmitted into the subsurface through a transmitting antenna and the resulting signal is received by a second antenna. The received signal may include an air wave, a ground wave, and reflected signals from subsurface boundaries, all in superposition.

The particular frequency range at which the radar operates is determined by radio frequency electronics and the specific antennas used. Typically in geologic materials, the lower the frequency range of the radar the greater the penetration range, assuming the transmitter output power and receiver sensitivity are not varied. The ability to resolve variations in electrical properties which have small spatial extent increases as the frequency increases, assuming a constant center-frequency to system-bandwidth ratio. It is necessary, therefore, to use antennas having the optimum frequency range and bandwidth characteristics to see the desired electrical variations in the particular geologic medium.

A GEODAR-I, Model-2441, or equivalent unit, will be used at the Carswell sites. A typical radar unit consists of a timing control unit which synchronizes all timing for the transmitter, receiver, data recorder, and data display. The transmitter and receiver electronics are located in the respective antenna modules. They are connected to the control unit through cables. Only the timing signals, the audio frequency facsimile of the received signal, and the DC voltage are carried on this cable.

The receiver has a sampling head similar to a sampling oscilloscope. The sampling head uses the repetitive nature of the received signal to transform the radio frequency signal into audio frequency facsimile. The transmitter sends out pulses and the receiver samples the received pulse for a small interval of the pulse duration. After each transmitter pulse, the sample window is moved in time and this is repeated until a complete scan of the pulse is obtained. The rapid rate at which the data is collected presents a virtually continuous record. During the survey, the antennas can be moved continuously over the line of interest. sampled audio frequency signal is sent to the control unit where it is amplified, filtered and then monitored on an oscilloscope. signal is recorded on an electrostatic recorder.

The electrostatic recorder uses a variable gray scale to display the data. Each scan draws a line across the paper in the direction of increasing signal travel time with the intensity dependent upon the radar signal amplitude.

The received radar signals are filtered before recording using audio frequency analog filters located in the control unit. Analog filtering helps to remove some of the equipment-generated noise.

For a routine reconnaissance map of reflections in the ground, the antennas are mounted rigidly at a known separation and moved along the profile line. The resultant trace shows reflection travel time versus position along the profile. In the profile mode, the travel time is related to the reflector depth and signal propagation velocity.

2.1.3 Geochemical Survey

A ground-water field screening will be performed at the POL Tank Farm.

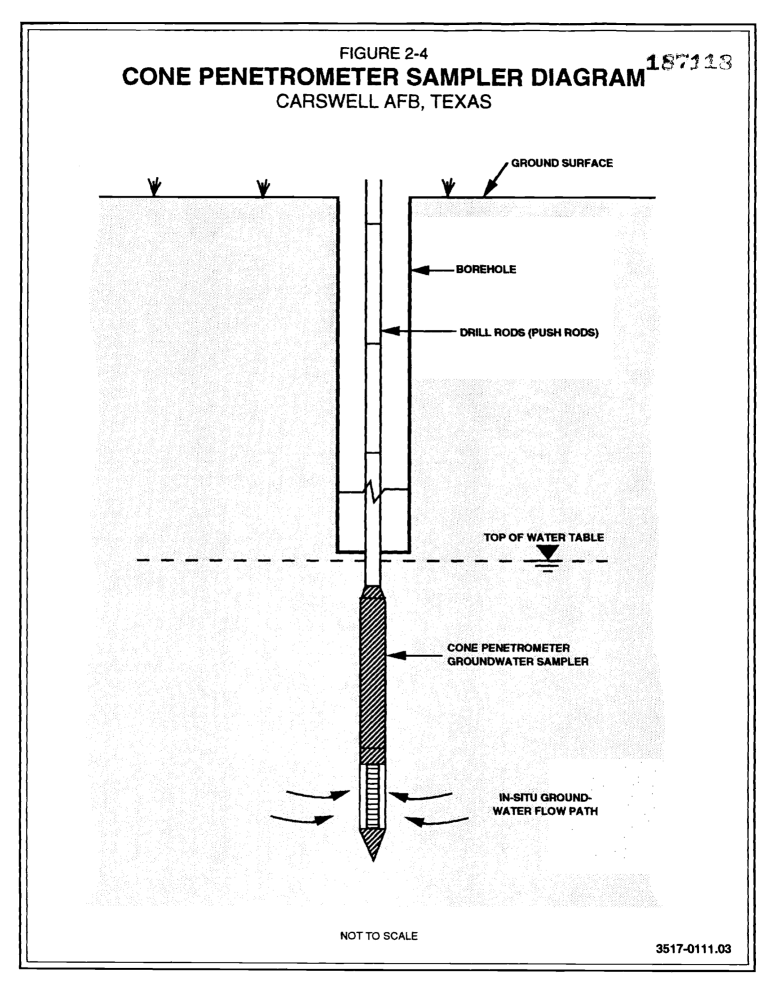
Ground-water samples will be collected for screening and analyses using a cone penetrometer type sampling system at Carswell AFB as part of the RCRA facility investigation effort. The intent of this activity is to conduct an evaluation of the nature and extent of ground-water contamination at Carswell AFB. The contaminants of concern are petroleum hydrocarbons. This screening and analyses will be conducted by Transglobal Environmental Geochemistry (TEG), Texas located in Austin, Texas. TEG utilizes hydraulic drivepoint rigs to perform the ground-water screening. The Strataprobe™ is a cone penetrometer type sampling system developed by TEG.

2.1.3.1 <u>Strataprobe™</u> - The Strataprobe™ ground-water field screening will be performed at the POL Tank Farm. The POL Tank Farm is located in the east area of Carswell AFB, west of and adjacent to Knight's Lake Road and north of Hobby Shop Road (Figure 2-1). Twenty-two Strataprobe™ samples will be collected at the POL Tank Farm to provide supplemental geochemical information in order to delineate the extent of the ground-water contamination. Three Strataprobe™ locations will be held in reserve for discretionary reasons once the ground-water field screening is implemented.

The sampling tool of the Strataprobe™ will be hydraulically pushed to the water table, which is estimated to be at 10 feet. Collection of the ground-water samples will be performed with a cone penetrometer equipped with a screened intake and sample chamber (Figure 2-4). The tool is advanced to the target depth and the outside drive casing is retracted 1 foot to expose the screen. Upon filling of the sample chamber, the tool can be withdrawn from the boring, and the sample transferred to the appropriate containers. These water samples will be analyzed in the field by TEG for the parameters listed in Table 2-5. Approximately 10 percent of these samples will be sent to an off-site laboratory for confirmation. The cone penetrometer will be decontaminated after each sample is containerized. Equipment decontamination procedures are described in Section 2.1.8.

2.1.4 Borehole Construction, Lithologic Sampling and Logging

A soil boring will be performed to allow installation of each monitoring well. The soil boring will provide subsurface information for well design and site stratigraphy. The soil boring will be observed by a qualified geologist or geotechnical engineer. The geologist/engineer will log the subsurface conditions encountered in the boring, and record the information on a soil



boring log. An example of a soil boring log is shown in Figure 2-5. Any other additional information concerning the boring will also be recorded on the log. Soils will be classified using the Unified Soil Classification System (ASTM D 2488-69).

Hollow-stem augering will be the preferred technique for all soil borings. The augers will have a minimum inner diameter (I.D.) of 6.25 inches to allow installation of the wells as described in subsequent report sections. It is anticipated that use of the hollow-stem augers will provide a sufficiently stable hole for soil sampling and monitoring well installation. If drilling fluids are to be utilized, a sample of the fluids introduced to the borehole will be analyzed to evaluate potential constituents introduced into the monitoring well.

2.1.5 Monitoring Well Construction and Installation

2.1.5.1 <u>Construction</u> - The monitoring wells will be installed to accomplish the following objectives: to collect representative ground-water samples; to prevent contamination of the aquifer by the drilling equipment; to prevent inter-aquifer contamination; and to prevent vertical seepage of surface water into the monitoring well water-intake zone. The equipment, procedures and personnel that will be used at the Carswell AFB to accomplish these objectives are discussed below.

The monitoring wells will be designed to intercept the upper portion of the water-table aquifer. The screen will be placed so that it intercepts the maximum upper fluctuation of the water table. Vertical fluctuation of the surficial aquifer water table at Carswell AFB generally does not exceed 5 feet over the course of a 12-month period (personal communication, J. Bartelino, U.S. Geological Survey, 1993). Seasonal fluctuations in the water table

FIGURE 2-5

187120

	SO	IL T	TES	T	BC	RI	NG	iF	RECORD	
J [V	JOB NO JOB NAME DATE: WEATHER DRILLER	_							BORING NO. G.S. ELEV. HOURS MOVING HOURS DRILLING PAGEOF	
DEPTH (FEET)	STRATA DESCRIPTION	#	1	2	3	4	N	P I D	R SOIL CLASSIFICATON AND REMARKS	DEPTH (FEET)
1										
-										-
			-							
-										==
_										
		-	\vdash							
-		F			_					
		F	-							
					F		F			
-										
_										
Ì			\vdash							
			-		\vdash		_	-		
BORING	G TERMINATED: G REFUSAL:					<u>-</u>	<u> </u>		HOD OF ADVANCING BORING	DEPTH
WATER	TOB DEPTH					-	Н.	AND	R AUGER CHOP: W/MUD: W/WATER	TO
WATER CASING	LOSSESLENGTH_					<u> </u>			RY DRILL: W/MUD: W/WATER OND CORE	TO TO
	C INSTALLED BY:CH		ED B	Y : _			DIS	CRE	PANCIES:	

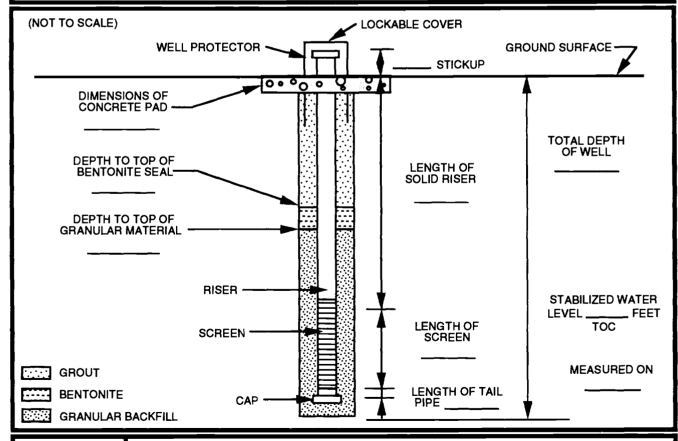
will be taken into account in placing the screen for each well. The screen will extend a minimum of 1 foot and a maximum of 5 feet above the water level encountered during drilling.

The screen and riser pipe will be installed through the hollow stem The pipe will be centered and suspended prior to placing the filter pack. Sand for the filter pack will be tremied into the annular space between the well casing and hollow stem augers using a tremie pipe. The hollow stem auger extensions will be withdrawn as the sand is placed into the well. The sand will be placed to a minimum of 2 feet above the top of the screen. If wells are installed so that the top of screen is at a depth of 2 feet or less, the sand pack will be installed to 6 inches above the screen. The sand will be continuously tamped to prevent bridging. depth of the sand pack and the amount of sand used will be continuously monitored. After the filter pack has been installed to a level of at least 2 feet above the top of the screen, the well will be surged for 10 minutes using a surge block. Additional sand will be added to bring the level back up to 2 feet above the screen. The well will then be surged for an additional 5 minutes. More sand will be added if necessary to return the level of the sand to 2 feet above the screen. Following placement of the sand, 3 feet of bentonite pellets will be placed above the sand pack. The remainder of the open hole will be sealed with a cementbentonite grout. The grout will be placed using a tremie pipe equipped with a side discharge. A Type II Well Completion Diagram is shown in Figure 2-6.

2.1.5.2 <u>Materials and Installation</u> - All monitoring well surface casing and riser pipe will consist of new, threaded, and flush joint, Schedule 40 polyvinylchloride (PVC). The pipe will conform to ASTM F-480-88A standards. The pipe will bear markings identifying the material as that which is specified. Each section

MONITORING WELL INSTALLATION DIAGRAM

BENTONITE TYPE GROUND SURFACE ELEVATION_____ MANUFACTURER _ CEMENT TYPE TOP OF SCREEN ELEVATION_ MANUFACTURER . REFERENCE POINT ELEVATION.... BOREHOLE DIAMETER ___ TYPE SAND PACK_ __ GRADATION__ SCREEN DIAMETER ______ SLOT SIZE__ SAND PACK MANUFACTURER_ LAW ENVIRONMENTAL, INC. SCREEN MATERIAL. FIELD REPRESENTATIVE _ MANUFACTURER_ DRILLING CONTRACTOR __ RISER MATERIAL AMOUNT BENTONITE USED___ MANUFACTURER_ AMOUNT CEMENT USED___ RISER DIAMETER _ AMOUNT SAND USED. DRILLING TECHNIQUE _ AUGER SIZE AND TYPE _____ STATIC WATER DEPTH (after dev.) ___ REMARKS _



QA/QC

INSTALLED BY: _____ INSTALLATION OBSERVED BY: ______
DISCREPANCIES: _____

will be joined by threaded flush-joint couplings to form watertight seals. No organic solvents or glue will be used in joining the pipe.

All well screens will consist of new, commercially fabricated, threaded 10-foot, flush joint, 4-inch PVC, factory slotted; slot size .010. A threaded PVC plug or short sand sump section will be provided for the bottom of the well.

A sand pack will be installed in the annulus between the boring and the well screen. The sand filter pack will extend 2 feet above the top of the screen. The sand pack will consist of clean, inert, non-carbonate materials. A "20/40" sand, or its equivalent, will be used in the wells to be installed at the sites.

A 3-foot thick bentonite seal will be placed in the annulus above the filter pack and will be allowed to hydrate to prevent intrusion of the grout into the filter sand. A cement grout will be placed in the annular space between the well casing and boring from the top of the bentonite seal to the ground surface. The cement mixture will consist of Portland cement (ASTM-C150), and water added in the proportion of no less than 5 to no more than 7 gallons per 94-pound bag of cement. Additionally, 3 percent by weight of bentonite powder will be added to the mixture to help reduce shrinkage, if permitted by state regulations.

Upon completion of the well, a vented cap will be installed to prevent material from entering the well. The PVC riser will be surrounded by a larger diameter steel casing rising approximately 24 to 36 inches above ground level. The steel casing will be provided with a locking cap and lock. A 3-foot square, 4-inch thick concrete pad, sloped away from the well, will be constructed around the well casing at the final ground level elevation. Three steel posts, with 3-inch diameters and 5-foot lengths, will be

equally spaced around the well outside of the concrete pad. Monitoring wells will be identified on the outside of the casing.

All wells will be checked for plumbness after installation. A 10-foot length of pipe, a half-inch less in diameter than the I.D of the riser pipe, will be run through the entire length of the well. This pipe section will be steam cleaned prior to use at each location. If the pipe does not run freely through the full length of the well, the well will be replaced or repaired.

- 2.1.5.3 <u>Well Development</u> The monitoring wells will be developed no sooner than 48 hours after grouting is completed. Development protocol will be as follows:
 - a. Measure static water level. Water levels will be measured using an interface probe capable of detecting both petroleum product and water levels.
 - b. Measure total well depth.
 - Develop the well using a combination of surging, bailing c. and pumping. Begin by surging the well with a surge block for a period of not less than 15 minutes. Then use either a bailer or a pump to evacuate the well. Continue pumping/bailing and periodically surging until: 1) the suspended sediment content of the water is less than 0.75 mL/L as measured in an Imhoff Cone according to Method E160.5; 2) the turbidity remains within a nephelometric turbidity unit (NTU) range for at least 30 minutes; and 3) the temperature, pH and conductivity have stabilized. Stabilization criteria are as follows: temperature plus or minus 1 degree celsius, pH plus or minus 0.1 units, conductance plus or minus 5 percent.

- d. All fluids introduced into the well will be removed during development. Therefore in addition to the above criteria at least three times the total quantity of fluids added during drilling will be removed during development.
- e. In the event that the well goes dry during development, the following steps will be used:
 - 1) Immediately after the well is purged dry, check the static water level and note the time of the reading. Suspend the probe of the static water level indicator 6 inches above the last static water level reading and record the time it takes the monitoring well to recharge to that level 6 inches above the last reading. Note the time again.
 - 2) Repeat this procedure over the next 6-inch interval to verify accuracy.
 - If the recharge rate is sufficient to recover to at least 80 percent of the original static water level in 1 hour, purge the well dry two additional times. Development will be completed after three purged dry cycles.
 - 4) If the recharge rate is not sufficient to recover the 80 percent of the original static water level in 1 hour, notify the Law Project Manager for further guidance.
- f. Denote physical characteristics of water throughout well development (color, odor, turbidity, etc.).

- g. Record the total quantity of water removed.
- h. Measure static water level after at least 24 hours.
- i. Measure total well depth.
- j. After final development collect approximately 1 liter of water from the well in a clear glass jar. Label and photograph the water jar. The photo will be suitably backlit to show the clarity of the water.

Well development data will be recorded on Monitoring Well Development Sheet, Figure 2-7.

2.1.6 Aguifer Tests (Slug Tests)

In-situ hydraulic conductivity tests will be performed on the new monitoring wells a minimum of 24 hours after ground-water sampling. Decontamination of downhole testing equipment will be performed in accordance with Section 2.1.8. The tests will be conducted to determine formation permeability. The results of the tests will be used to estimate ground-water flow rates.

The tests which will be performed are known as rising head tests. The rising head test involves removing a solid rod which was previously inserted into the water column in the well. The rising head test is performed once the water level has returned to static after the initial insertion of the rod. Removal of the rod causes an immediate lowering of water level. The water level recovery to static is recorded over time using an automatic data logger. Water level readings will be measured and recorded continuously until the recovery is 90 percent of the original static water level. The rate of recovery is controlled by formation characteristics.

JOB NAME			
BY	CHECKED	SHEET	OF
	WELL DEVELOPM	ENT DATA	
1. Well No			
2. Date of Installation :			
Date of Development :			
Static Water Level : Before Devel	-	ft.: 24 Hours After	ft
5. Quantity of Water Loss During Dril	-		
6. Quantity of Standing Water in Well	_		Gal.
, ,	Start	During	<u>End</u>
7. Specific Conductance (umhos/cm			
Temperature (cº)			
pH (s.u.)			
8. Depth From Top of Well Casing to	Bottom of Well	ft. (from Well Ir	nstallation Diagram)
9. Well Diameter			
10. Screen Length	ft.		
11. Minimum Quantity of Water to be	Removed (5 Well Volumes)	
12. Depth to Top of Sediment: Befo	re Development	ft.; After Developme	ent ft.
13. Physical Character of Water:			
14. Type and Size of Well Developmen	nt Equipment :	. <u>-</u>	
15. Description of Surge Technique, If	Used :		
16. Height of Well Casing Above Grou	nd Surface :	tt. (from Well Instal	lation Diagram)
17. Quantity of Water Removed :	Gal.	Time for Removal :	Hr./Min
18. 1-Liter Water Sample Collected : _	(Tin	ne)	
19. Turbidity in Nephelometric Units _	NT	Us	

^{*}Development Conditions: 1) Well Water if Reasonably Clear
2) Sediment Thickness 5% of Screen Length
3) Removal of 5 Well Volumes, Including Saturated Filter Annulus
4) Stabilization of Specific Conductance and Water Temperature

FIGURE 2-7 (CONT'D)

WELL DEVELOPMENT DATA

JOB NAME		JOB No	_		_	
BY	CHECKED		SHEET	2	OF	2
Well No						

Date/ Time	Hrs Dev/ Cum Hrs Dev	Gals Purged/ Cum Gals Purged	рН	Temp	Cond	Remarks
				,	!	
					_	
		_		:		
					!	

The data results of the hydraulic conductivity test are then The following formula (Bouwer, 1989) is utilized to calculated. calculate hydraulic conductivity (K):

K (ft/sec) =
$$\frac{r_c^2 \ln (R_e/r_w)}{2 L_e} * \frac{1}{t} * \frac{\ln Y_0}{Y_t}$$

where:

 r_2 (ft) = well radius

= effective radial distance over which R_a (ft)

the head difference is dissipated

radial distance between well center r_u (ft) =

and undisturbed aquifer

 L_{p} (ft) = height of saturated screen

 Y_0 (ft) = water level Y at time zero Y_+ (ft) = water level Y at time t

Y, (ft)

 $t (sec) = time since Y_0$

Analysis of the data will be performed utilizing a commerciallyavailable computer program such as AQTESOLV®. Because we expect the screen will extend above the water table, only the rising head data will be used to calculate K.

2.1.7 Surveying

Upon completion of the wells, a state of Texas certified surveying crew will locate by standard surveying methods each new monitoring well and sampling location. A vertical survey will be conducted to establish the elevation of each permanent monitoring well. Vertical control will be to the National Geodetic Vertical Datum. The horizontal grid coordination of each monitoring well to within 0.1 foot and the ground elevation to within 0.01 foot and the elevation for each well within 0.01 foot will be recorded. The survey will be tied to the state plane system.

2.1.8 Equipment Decontamination

The following decontamination procedures will apply to split spoons, hand augers, stainless-steel bowls and spoons, and ground-water bailers. The decontamination steps are as follows:

- 1) Hand wash with bristle brush and a solution of Alconox (or equivalent).
- 2) Rinse with copious quantities of potable water.
- 3) Rinse with deionized (Reagent Grade II) water.
- 4) Spray-rinse with pesticide grade methanol.
- 5) Spray-rinse with pesticide grade hexane.
- 6) Air dry.

The following procedure shall be used to decontaminate large pieces of equipment such as drill rigs, auger flights, and casing.

- Wash the external surface of equipment or materials with high-pressure hot water and Alconox or equivalent, and scrub with brushes if necessary until all visible dirt, grime, grease, oil, loose paint, rust flakes, etc., have been rinsed from the equipment.
- 2) Rinse with potable water.
- Air dry. The sampling equipment will be cleaned prior to each use in accordance with this procedure. All decontamination solutions will be stored on site until analyses have been completed.

The following procedure will apply to the decontamination procedure for the Strataprobe $^{\text{\tiny{M}}}$ sampling equipment used for the geochemical survey:

- 1) Remove all loose dirt.
- 2) Wash with Alconox.
- 3) Rinse with potable water.
- 4) Rinse with deionized (Reagent Grade II) water.

2.1.9 Waste Handling

All drill cuttings, drilling fluid and other investigation derived wastes (IDW) will be containerized in clean, Department of Transportation (DOT), 17 H drums. All drums of IDW will be adequately labeled and moved to a staging area designated by the base. Each drum of IDW will be marked at the start of waste accumulation with the following information documented on the drum:

- · Date upon which accumulation begins
- · Site identification
- · Boring identification number
- · Drum contents (soil, wash, water, etc.)

Upon completion of the field program, a drum log (inventory) will be prepared that accounts for all drums and wastes generated drums site activities.

Law will analyze a maximum of one composite sample from the drums for each area to be analyzed for TCLP. The analytical information will be used to prepare a manifest for ultimate shipment of the drums to a disposal facility. Law will be responsible for off-site disposal of any drums.

Development water will be drummed and moved to a staging area on base following the procedure outlined in the drill cutting disposal section above. Drummed development water will not be analyzed, but the results of water analysis after monitoring well sampling will be used to prepare a manifest for ultimate shipment of the drums to a disposal facility. Law will be responsible for off-site disposal of any drums.

A separate drum will be provided for additional wastes, such as gloves, tyvek, etc., generated in drilling and sampling operations, which will also be included as IDW to be disposed of along with the other drums of IDW.

2.1.10 Field Activity Summary

The field tasks and field samples per site are presented in Table 2-1.

2.2 <u>ENVIRONMENTAL SAMPLING</u>

Field sampling activities include the collection of ground-water, surface water, sediment, and soil samples. The collection methods, including sample handling, sample custody, QC samples, and sample analysis are presented in the following subsections. Table 2-2 provides a list of equipment required.

2.2.1 Procedures for Collection of Samples

This section presents the planned program for collection of samples for chemical analysis. The types and numbers of field quality control samples are presented in Tables 2-3 through 2-7 by matrix and parameter. The following sections present the site-specific

TABLE 2-1

FIELD TASKS RCRA Facility Investigation Carswell Air Force Base, Texas

SITE		FIELD TASKS		RATIONALE
POL TANK FARM	- -	Perform geophysical survey	1.	Determine the location of any pipechase or utilities.
	2.	Perform ground-water scree geochemical survey using Spush method utilizing a cortwenty five upper zone grouscreen for contamination.	2. 0	ining. Perform 2. Determine the extent of ground-water contamination from fuel and expand penetrometer. Collect ground-water data base.
UNNAMED STREAM	.	Perform geophysical survey	1. L	Determine locations of USTs, piping and utilities.
	2	Install three new monitoring wells (MW-01 through MW-03). Screen soils with OVA/PID. Collect two soils samples per boring for lab analyses, one from the zone of the highest OVA/PID reading and one from the ground-water interface. Two additional soil samples will be collected from the upgradient soil boring.	2. 8	Assess the hydrocarbon impact to native soils.
	က်	Sample ground water on the three new monitoring wells and sample six existing wells for a total of three rounds of ground-water sampling.	3. 1. 1.	Determine the nature and extent of ground-water contamination and expand the ground-water data base.
	4.	Surface water/sediment sampling.	4.	Determine the extent of contamination related to the oil/water separator
	5.	Aquifer Testing (In-Situ hydraulic conductivity)	.5	Determine hydraulic characteristics of the upper zone aquifer.

TABLE 2-2

FIELD EQUIPMENT CHECKLIST Remedial Investigation/Feasibility Study Carswell Air Force Base, Texas

187134

<u>General</u>	
1.	Investigation Work Plan
2.	Sampling and Analysis Plan
3.	Health and Safety Plan
4.	Site base map
<u> </u>	Hand calculator
6.	
	Personal clothing and equipment
Cafety Favinas	
Safety Equipm	
1.	Work gloves
2.	Work gloves Barrier rope Plastic flagging tape
3.	Plastic flagging tape
4.	Drinking water container
5.	Gatorade or equivalent
6.	First aid kit
7.	
8.	Spill control (Vermiculite)
9.	Knife Flashlight or lantern
10.	Flashlight or lantern
	Personal locator beacon (sound, radio or light)
	Radio, two-way
13.	Safety belt, harness, and lifeline
Personal Prote	ctive Equipment
1.	Long cotton underwear
2.	Coveralls, cotton
3.	Coveralls, Tyvek
4.	Surgical gloves
5.	Protective gloves (polypropylene/nitrile/viton)
6.	
7.	Hard hat
8.	Face shield
9.	Safety glasses
10.	Air-purifying respirator
11.	Respirator spectacle kit

FIELD EQUIPMENT CHECKLIST Remedial Investigation/Feasibility Study Carswell Air Force Base, Texas

Sampling Equi	pment
1.	Tool box with assorted tools (pipe wrenches, screwdrivers, socket set and driver, open and box end wrenches, hacksaw, hammer, vise grips)
2.	Geologic hammer
3.	Trowel
4.	Stainless steel and/or Teflon spatula
5.	Hand auger
6.	Dames & Moore Type U split-spoon sampler with liners (stainless steel, brass)
7.	Engineer's tape
8.	Steel tape
9.	Electric water level sounder
10.	Petroleum interface probe
11.	Batteries
12.	Bailers (Teflon, stainless steel, acrylic, PVC)
13.	Slug test water displacement tube
14.	Vacuum hand pump
15.	Electric vacuum pump
16.	Displacement hand pump
17.	Mechanical pump (centrifugal, submersible, bladder)
18.	Portable generator
19.	Gasoline for generator
20.	Hose
21.	Calibrated buckets
22.	Stop watch
23.	Orifice plate or equivalent flow meter
24.	Data logger and pressure transducers
25.	Sample bottles
26.	Sample preservatives (nitric, hydrochloric, sodium hydroxide)
27.	Heavy-duty aluminum foil
28.	Coolers
29.	Ice (not reusable ice packs ["Blue Ice"])
30.	Large "Ziploc" freezer bags
31.	Miscellaneous garbage bags
32.	Duct tape
33.	Strapping tape
34.	Paper towels
35.	"Bubble" pack
36.	Vermiculite
37.	Clear tape
38.	Containers for field tests (pH, SC, temp.)
39.	Transfer pipets (glass) and bulbs
40.	Polyethylene rope for bailers
41.	Teflon tubing to cover bailer rope

FIELD EQUIPMENT CHECKLIST Remedial Investigation/Feasibility Study Carswell Air Force Base, Texas

187136

1. Shovels 2. Keys to well caps 3. pH meter (with calibration solutions) 4. pH paper 5. Thermometer 6. Conductivity meter (with calibrating solution) 7. Organic vapor analyzer or photoionization detector with calibration gas 8. Explosimeter
2. Keys to well caps 3 pH meter (with calibration solutions)
3 pH meter (with calibration solutions)
4. pH paper 5. Thermometer 6. Conductivity meter (with calibrating solution) 7. Organic vapor analyzer or photoionization detector with calibration gas
5. Thermometer 6. Conductivity meter (with calibrating solution) 7. Organic vapor analyzer or photoionization detector with calibration gas
6. Conductivity meter (with calibrating solution) 7. Organic vapor analyzer or photoionization detector with calibration gas
7. Organic vapor analyzer or photoionization detector with calibration gas
8. Explosimeter
9. Oxygen meter
10. Turbidimeter
Decontamination Equipment
1. Potable water
2. Alconox, Liquinox, or equivalent
3. Methanol, pesticide-grade
4. Hexane, pesticide-grade
5. Nitric acid
6. Distilled, deionized water, Type II reagent grade
7. Teflon squeeze bottles
8. Stainless steel garden sprayers
8. Stainless steel garden sprayers Buckets and containers for decontamination fluids
10. Scrub and bottle brushes
11. Waste containers (plastic trash cans/55-gallon hazardous waste containment drums)
12. Steam cleaner (generally provided by subcontractor)
Documentation Supplies
1. Field log book
2. Daily quality control report
3. Soil test boring record
4. Monitoring well installation log forms
5. Well development data forms
6. Aquifer test data forms
7. Sample chain-of-custody forms
8. Custody seals
9. Cooler labels ("This side up," "Hazardous Material," "Fragile")
10. Courier labels/airbills
11. Request for analysis forms
12. Sample bottle labels
13. Camera and film
14. Paper
15. Pens/pencils
16. Indelible ink pens

TABLE 2-3

SAMPLING SUMMARY: SUB-SURFACE SOIL FROM BORINGS RCRA Facility Investigation - Carswell Air Force Base, Texas

Volatile Organics (SW8240) 2 2 4 1 4 4 4 1 1 1 4 <th>PARAMETER</th> <th>TOTAL NO. FIELD LOCATIONS</th> <th>SAMPLES PER LOCATION</th> <th>TOTAL NO. FIELD SAMPLES</th> <th>Field E Duplicate</th> <th>FIELD QC Equipment Blank</th> <th>FIELD QC SAMPLES quipment Ambient Blank Blank</th> <th>Trip Blank (a)</th> <th>TOTAL NO. LAB SAMPLES</th>	PARAMETER	TOTAL NO. FIELD LOCATIONS	SAMPLES PER LOCATION	TOTAL NO. FIELD SAMPLES	Field E Duplicate	FIELD QC Equipment Blank	FIELD QC SAMPLES quipment Ambient Blank Blank	Trip Blank (a)	TOTAL NO. LAB SAMPLES
S 3 6 8 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	(SW8240) am: Downgradient (b) am: Upgradient (c)		2 4 9	4 4 80	2	0	a	2	8 8 16
S 3 6 8 2 2 0 0 1 1 1 0 0 0 1 1 1 1 1 1 1 1 1 1	(SW8020) m: Downgradient m: Upgradient		0 4 0	4 4 8	0	2	a		8 8
3 1 3 1 1 0 0 0 S 3 1 1 1 0 0	Total Recoverable Petroleum Hydrocarbons (E- Unnamed Stream: Downgradient Unnamed Stream: Upgradient		0 4 0	4 4 8	a	6	0 0	0 0 0	6 6 21
	Toxicity Characteristic Leaching Procedure (d) Unnamed Stream (SW1311) TOTALS		- -	က က			0 0	0 0	ហ

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan.

(a) Volatile organics only; number to be initiated will depend upon number of shipments.
(b) Soil samples from oil/water separator and unnamed stream monitoring well installations.
(c) Soil samples from single upgradient monitoring well installation.
(d) TCLP analyses will be performed on the containerized waste material generated from the drill cuttings.

SAMPLING SUMMARY: GROUND WATER — MONITORING WELLS RCRA Facility Investigation — Carswell Air Force Base, Texas

		TOTAL NO.	TOTAL NO.	TOTAL NO.		FIELD OC SAMPI ES	AMPLES		TOTAL
PARAMETER		FIELD LOCATIONS	SAMPLING EPISODES	FIELD SAMPLES	Field Duplicate	Equipment Blank	Ambient Blank	Trip Blank (a)	LAB SAMPLES
Volatile Organics (SW 8240) Unnamed Stream		6	თ	27	ო	က	က	က	39
	TOTALS	6	8	27	က	က	က	ဇ	39
Total Recoverable Petroleum Hydrocarbons (E418.1) Unnamed Stream		6	ო	27	ო	ო	0	0	33
	TOTALS	6	က	27	ဇ	ဇ	0	0	33
Oil and Grease (SW 9071) Unnamed Stream		თ	ო	27	ო	ო	0	0	33
	TOTALS	6	က	27	က	က	0	0	33
		თ	ო	27	ო	ო	0	0	ဗ
Unnamed Stream: Dissolved		6	က	27	က	ო	0	0	33
	TOTALS	18	9	54	9	9	0	0	99
Arsenic (SW 7060) Unnamed Stream: Total		6	ო	27	ო	ო	0	o	33
Unnamed Stream: Dissolved		6	က	27	ဇာ	က	0	0	33
	TOTALS	18	9	54	9	9	0	0	99
								18	
3517-0111.10								7138	1 of 2

TABLE 2-4

SAMPLING SUMMARY: GROUND WATER - MONITORING WELLS RCRA Facility Investigation - Carswell Air Force Base, Texas

ed TOTALS 18 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	PARAMETER		TOTAL NO. FIELD	TOTAL NO. SAMPLING	TOTAL NO. FIELD	Field	FIELD QC SAMPLES Equipment Ambient	SAMPLES	Trip	TOTAL NO. LAB
TOTALS 18 9 9 10 10 10 10 10 10 10 10 10 10 10 10 10	= =		തത	တ က	27	3 3	o s	Blank 0 0	Blank (a) 0 0	SAMPLES 33 33
d TOTALS 18 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9		TOTALS	18	9	54	9	9	0	0	99
TOTALS 18 9 TOTALS 18 5			တတ	თ თ	27 27	ღ ღ	ოო	00	0 0	33 33
9 TOTALS 18		TOTALS	18	9	54	9	9	0	0	99
TOTALS 18 5 TOTALS 5	m: Total m: Dissolved		တတ	ო ო	27 27	ო ო	ო ო	0 0	0 0	33 33
TOTALS		TOTALS	18	9	54	9	9	0	0	99
			ß	-	ហ	-	-	0	0	~
	-	rotals	2		2		-	0	0	7

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan.

(a) Volatile organics only; number to be initiated will depend upon number of shipments. (b) The purge water will be analyzed for the parameters of conern in the TCLP list.

TABLE 2-5

SAMPLING SUMMARY: GEOCHEMICAL SURVEY GROUND WATER - FIELD SCREENING RCRA Facility Investigation - Carswell Air Force Base, Texas

		TOTAL NO.	TOTAL NO.	FIELD	FIELD QC SAMPLES	ES		OFF-SITE	TOTAL NO.
(SW 8020) 9 9 1 1 1 1 1 TOTALS 10 10 2 2 2 1 3 3 1 1 1 0 TOTALS 25 25 3 3 0 TOTALS 10 10 1 1 0	PARAMETER	FIELD LOCATIONS	FIELD SAMPLES	!	Equipment Blank	Ambient Blank	Trip Blank(a)	CONFIRM. ANALYSES (b)	LAB SAMPLES
TOTALS 10 10 2 2 2 1 1 1 0 0 10 10 10 10 10 10 10 10 10 10	Aromatic Volatiles (SW 8090)								
TOTALS 10 10 2 2 1 1	POL Tank Farm	6	6	-	-	-	-	-	13
TOTALS 10 10 2 2 1 1	Reserve (c)	-	-	-	-	0	0	0	ო
TOTALS 10 10 10 10 10 10 10 10 10 10 10 10 10	TOTALS	10	10	2	2	-	-	-	16
TOTALS 25 25 3 3 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Petroleum Hydrocarbons (SW 8015M)	66	8	٥	0	o	0	ო	59
TOTALS 25 25 3 3 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Reserve (c)	່ພ	က	ı 	· -	0	0	0	ഹ
9 9 1 1 0 0 1 0 0 0 0 0 1 1 1 0 0 1 0 1	TOTALS	25	25	က	က	0	0	ဇာ	34
TOTALS 10 10 1 0 0 0 0 0 0 0 0	Lead (SW 7421) POL Tank Farm	6	ნ	-	-	0	0	0	1
10 1 1 0	Reserve (c)	-	-	0	0	0	0	0	-
	TOTALS	10	10		-	0	0	0	12

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan.

(a) Volatile Organics only; number to be initiated will depend upon number of shipments.
 (b) Methods SW 8020 and SW 8015M will have ten percent of all samples sent to an off-site laboratory for confirmatory analyses.
 (c) Three ground-water sampling locations will be reserved for discretionary use during the implementation of the field work.

TABLE 2-6

SAMPLING SUMMARY: SURFACE WATER RCRA Facility Investigation — Carswell Air Force Base, Texas

PARAMETER		TOTAL NO. FIELD LOCATIONS	TOTAL NO. FIELD SAMPLES	Field Duplicate	FIELD QC SAMPLES Equipment Ambient Blank Blank	SAMPLES Ambient Blank	Trip Blank (a)	TOTAL NO. LAB SAMPLES
Volatile Organics (SW 8240) Unnamed Stream		ო	ო	-	-	-	-	7
	TOTALS	ဇ	က		-	-	_	
Total Recoverable Petroleum Hydrocarbons (E418.1 Unnamed Stream	rbons (E418	3	ო	-	-	0	0	Ŋ
	TOTALS	ဇ	ო		-	0	0	2
Oil and Grease (SW 9071) Unnamed Stream		ო	ო	-	-	0	0	വ
	TOTALS	က	က		-	0	0	2

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan. (a) Volatile organics only; number to be initiated will depend upon number of shipments.

TABLE 2-7

SAMPLING SUMMARY: SEDIMENT RCRA Facility Investigation — Carswell Air Force Base, Texas

TOTAL NO. LAB SAMPLES	ഹ ഹ
Trip Blank (a) §	0 0
l II	0 0
FIELD QC SAMPLES Equipment Ambient Blank Blank	- -
Field Duplicate	
TOTAL NO. FIELD SAMPLES	ო ო
TOTAL NO. FIELD LOCATIONS	က က
PARAMETER	ICP Screen for Metals (SW 3050/6010) Unnamed Stream TOTALS

Note: Analytical methodologies are presented in Section 1.8.1 of the Sampling and Analysis Plan. (a) Volatile organics only; number to be initiated will depend upon number of shipments. sampling requirements, the planned sample collection procedures, and the procedures to be used to maintain sample integrity.

Prior to sampling, field instruments will be calibrated, files containing sample information will be processed and labels will be prepared. Sample bottles will be sorted for each sample location according to analyses. Conditions and sampling information will be recorded in the field sampling books and used to assess sampling procedures in relation to the sample data. The field team leader will brief the sampling team on safety, decontamination stations and any other sampling protocols necessary. Each sampling team member will wear the appropriate level of safety gear as specified for each site in the Health and Safety Plan.

2.2.1.1 <u>Ground-Water Sampling</u> - Ground-water samples will be collected from newly installed wells. Ground-water samples will be obtained using cleaned and dedicated Teflon® bailers. Bailers will be decontaminated as outlined in Section 2.1.8. After the bailer has air-dried it will be wrapped in aluminum foil to prevent contamination. Polyethylene rope will be used to lower the bailer into the well. A Teflon® leader will precede the rope attached to the bailer to prevent the rope from entering the well during sampling.

Ground-water sampling will be conducted in a manner which minimizes interaction of the sample and the surface environment. The sampling protocol will be as follows:

a. <u>Floating Hydrocarbon Measurement</u> - An electronic interface probe will be used to test for the presence of floating product in the newly installed and existing wells. The probe will be lowered into the wellbore and if product is present, its thickness will be recorded.

Its presence will be confirmed by withdrawing a sample. The probe will be decontaminated after each use by the protocol described in Section 2.1.8.

b. Measure Water Level - An electronic water level indicator will be used to measure the static water level in the well prior to purging. The water level probe will be lowered into the wellbore and the water level will be recorded. The volume of water in the wellbore and filter pack will be calculated using the equation in subpart c.

Decontamination of the probe will be performed as per Section 2.1.8 to minimize the potential for crosscontamination between wells as the probe is being withdrawn from the well.

Purge the well until the pH, specific conductance, and c. temperature have stabilized within plus or minus 0.1 pH units, plus or minus 5 percent, and plus or minus 1 degree celsius, respectively. Remove a minimum of three fluid casing volumes, checking the above mentioned parameters after each casing volume is removed. after three fluid casing volumes have been removed, the parameters have not been stabilized, remove well casing volumes until the parameters stabilize or until six volumes have been purged. If the well is purged to dryness and does not recharge within a reasonable time to permit three volumes to be removed, record the volume removed and sample as soon as sufficient recharge has occurred. The calculation for well volume is presented below:

CALCULATING WATER VOLUME VOLUME OF WATER IN WELL

 H_1 π 7.48 [($R_1^2 - R_2^2$) . \emptyset + R_2^2]

Volume of Volume of PVC with filter pack filter pack

 H_1 = height of water in well screen and filter pack

in feet

 $\pi = 3.14159$

 R_1 = radius of borehole in feet

 R_2 = radius of PVC in feet

7.48 = amount of water in 1 cubic ft.

ø = porosity of filter pack

- d. <u>Label Sample</u> Label the sample container as outlined in Section 2.2.2.
- e. <u>Collect Sample</u> Ground-water sampling will be conducted by using dedicated bailers. The collection procedure is described below.

After purging the well and sufficient recharge has been allowed, samples for volatile organics should be collected using the first bailer volume. Lower the bailer slowly until it contacts the water surface, and allow the bailer to sink to the desired depth and fill, with a minimum of surface disturbance. Slowly withdraw the bailer, taking care to prevent contact of the bailer line with the ground. Slowly discharge the contents into the appropriate sample containers. Repeat the process as necessary to fill each container to the required volume (see Section 2.2.2.1). Vials for volatile analysis will

be completely filled, leaving no air space above the liquid portion (to minimize volatilization). Check that the Teflon on the Teflon®-lined silicone septum is toward the sample in the caps and secure the cap tightly. If semi-volatile compounds are to be sampled for, collect these samples next. Proceed to the collection of samples for the remaining analyses. Care will be taken for all pre-preserved bottles. If acids are present, the bottle will be opened downwind of and away from the body. All samples for dissolved metals should be collected last and filtered through a 0.45 μ m filter within 15 minutes after collection. A Millipore filtration system will be utilized for this procedure. The procedure for Millipore filtering is as follows:

- 1) Assemble the Millipore filtration unit. Assembly procedures can be found in Appendix G.
- 2) Using a funnel, pour test material to be filtered into the unit through the Tri-Clamp opening. Attach the TC hose adaptor (fitted with 1½-inch gasket) to the TC connection of top plate, using the stainless steel TC clamp provided.
- 3) Place appropriate receiving vessel below TC sanitary outlet or attach PVC hose to TC adaptor and run hose to appropriate container.
- 4) Slowly increase pressure on nitrogen tank. When liquid flow begins from outlet of assembled holder, immediately turn regulatory valve handle a few turns to lower pressure. Bleed air from top-plate vent valve slowly until pressure gauge reads between 10 and 15 psi. Close vent valve and (if necessary) readjust regulatory valve to set

pressure between 10 and 15 psi. Continue filtration until flow ceases.

- 5) Increase inlet pressure stepwise in 10 psi increments to 75 psi and continue filtration until flow ceases. No leaks should be apparent.
- 6) Shut off pressure from nitrogen tank regulator outlet valve, then open vent valve gradually to release pressure from cylinder.
- 7) Remove inlet TC hose connector and handwheel bolts, then remove top plate and extension cylinder of holder.
- 8) Acidify the sample with nitric acid to a pH of less than or equal to two. Test with pH paper.
- 9) Decontaminate the filtration apparatus as outlined in Section 2.3.4.
- f. <u>Measure pH. Temperature and Specific Conductance</u> -Follow procedure as outlined in Section 2.3.
- g. <u>Custody</u>, <u>Handling and Shipping</u> Complete the procedures as outlined in Sections 2.2.2 and 2.2.3.
- h. Measure Water Level After samples have been collected, the well cap should be replaced. Static water level will be measured again after sufficient recharge has been allowed (24 hours). This additional measurement of the water level will follow the procedure outlined in Section 2.2.1.1.b.

2.2.1.2 <u>Soil Sampling</u> - Soil samples will be collected from soil borings of the newly installed monitoring wells. The following section discusses soil sampling and collection procedures. Included are discussions of soil sampling protocol and equipment.

Subsurface soil samples from borings will be collected using a 3-inch, carbon-steel, split-barrel sampler with California brass ring inserts. Care has been taken to determine the best practical sampling procedure that will result in obtaining representative samples. The samples must maintain the integrity of the original medium through collection, transportation, and delivery to the analyst. The soil samples will be collected and containerized as described in the following paragraphs.

Subsurface soil samples from soil borings - The auger will encase an 24-inch long, carbon-steel split barrel sampler which will, in turn, encase four 6-inch California brass rings. Each boring will be advanced using hollow-stem augers. The sampling device will be placed on a sheet of aluminum foil upon retrieval from the auger. Each end of the sampling device will be opened by unscrewing the Samples from certain borings are to be collected at depths which exhibit the highest organic vapor readings. portable organic vapor analyzer (OVA) will be used to screen the individual boring samples. The sampling device will be opened and the end of each brass ring screened. After the boring samples have been screened for organic vapor, the brass rings will be sealed with a Teflon® patch and plastic cap, thus encasing the sample in the brass ring. The middle 6-inch brass ring sample will be sent to the laboratory for volatile organic compound analysis. event that soil recoveries are low, the selection of brass rings used for volatile analysis will be made on the basis of which rings contain 100 percent recovery to send to the laboratory. If none of the rings have 100 percent recovery, then the ring with the most soil will be capped and labeled, indicating it is less than 100 percent. Soil from the remaining two brass rings will be removed and placed into a stainless-steel mixing bowl. The sample will be thoroughly mixed with a stainless-steel spoon and placed into the appropriate pre-cleaned jars (see Section 2.2.2.1). All sampling equipment will be decontaminated as presented in Section 2.1.8.

- 2.2.1.3 <u>Surface Water Sampling</u> Surface water samples will be collected with stainless-steel beakers in "low-flow" areas to avoid the loss of volatile organics. Each beaker will be cleaned using the method described in Section 2.1.8 before sample collection. Collect samples from the downstream to the upstream locations. Slowly submerge the beaker into the water. When the beaker is full, slowly pour the contents down the side and into the sample bottles. Completely fill volatile vials first, leaving no headspace. Fill sample containers for the remaining analyses. Surface water samples should be collected from downstream to upstream and should be collected before sediment samples collected at the same location.
- 2.2.1.4 <u>Sediment Sampling</u> Sediment samples will be collected with stainless-steel hand augers. The sample will be obtained by driving the hand auger through the water and into the sediments. Rotate the auger as necessary to reach a sample depth of 6 inches, avoiding excess disturbance of surface sediment particles. Retrieve the auger by lifting upward and empty the contents of the auger bucket into a stainless-steel mixing bowl. Completely fill sample containers to be submitted for analysis with volatile parameters first, leaving no headspace. Once the volatile organic samples have been collected, mix the remaining sediment thoroughly and fill sample containers for the remaining analyses. Sediment samples should be collected from downstream to upstream and should be collected after surface water samples have been collected at the same location.

2.2.2 Sample Handling

The following sections describe the proper way to containerize, label, identify, ship, and preserve samples.

- 2.2.2.1 <u>Sample Containers, Amounts and Preservation</u> Samples will be placed in appropriate containers as required by AFCEE (AFCEE, 1991). The amounts to be collected and preservation procedures specified by AFCEE will be followed. These requirements are specified in Tables 2-8 (soil, sediment) and 2-9 (ground water, surface water).
- 2.2.2.2 <u>Sample Labels</u> Field sampling personnel must properly identify all samples taken in the field with an adhesive sample label attached to the sample container. The sample label must contain the site name, field identification number; the date, time, and location of sample collection; designation of the sample as a grab or composite; notation of the type of sample (e.g., groundwater, soil boring, etc.); identification of preservatives used; any remarks; and the initials of the sampler. The sample labels will be placed on the bottles so as not to obscure any QA/QC data, and sample information will be legibly printed with waterproof ink. Field identification must be sufficient to allow easy cross-reference with the site logbook. Clear tape will be placed over the label to prevent removal or damage.
- 2.2.2.3 <u>Sample Identification</u> The following section describes how each sample will be identified and what this identification signifies. Note that all sample IDs shall be no more than 10 characters long to facilitate input into the IRPIMS database.

For field samples, the following sample ID scheme shall be followed:

TABLE 2-8

SAMPLE CONTAINERS, AMOUNTS, AND PRESERVATION SOIL AND SEDIMENT SAMPLES Remedial Investigation/Feasibility Study Carswell Air Force Base, Texas

PARAMETER (a)	SAMPLE CONTAINER (b)	AMOUNT	PRESERVATION
Volatile Organics (SW 8240) (includes samples for Aromatic Volatiles analysis (SW 8020))	Two 2 oz. G septa jars	Full; no headspace	Cool to 4°C
Total Recoverable Petroleum Hydrocarbons (418.1)	One 8 oz. G jar (c)	Full	Cool to 4°C
Metals, total (SW 6010)	One 8 oz. G jar	Full	Cool to 4°C
Toxicity Characteristic (SW 1311) Leaching Procedure	One 8 oz. G jar (c)	Full	Cool to 4°C

NOTES:

- (a) Hold times are presented in Section 1.8
- (b) All containers to have Teflon®-lined lids except for vials for volatile organics which will have Teflon®-lined septa. Container codes: G = glass AG = amber glass P = polyethylene
- (c) Samples chosen for MS/MSD and/or Laboratory Duplicate require double the number of containers indicated.

TABLE 2-9

SAMPLE CONTAINERS, AMOUNTS, AND PRESERVATION-GROUND WATER AND SURFACE WATER SAMPLES Remedial Investigation/Feasibility Study

Carswell Air Force Base, Texas

	CAMPLE		
PARAMETER (a)	SAMPLE CONTAINER (b)	AMOUNT	PRESERVATION
Volatile Organics (SW 8240) (includes samples for Aromatic Volatiles analysis (SW 8020))	Three 40 mL G vials	Full; no headspace	HCl to pH < 2; Cool to 4°C
Petroleum Hydrocarbons (SW 8015M)	Three 40 mL G vials	Full; no headspace	H_2SO_4 to pH <2; Cool to $4^{\circ}C$
Total Recoverable Petroleum Hydrocarbons (418.1)	Two 1 L AG bottles(c)	7/8 full	H_2SO_4 to pH <2; Cool to $4^{\circ}C$
Metals, total & dissolved (SW 6010)	One 1 L P bottle	7/8 full	HNO ₃ to pH < 2; Cool to 4°C
Arsenic (SW 7060)	One 1 L P bottle (c)	7/8 full	HNO ₃ to pH < 2; Cool to 4°C
Mercury (SW 7471)	One 1 L P bottle	7/8 full	HNO_3 to pH < 2; Cool to $4^{\circ}C$
Selenium (SW 7740)	One 1 L P bottle	7/8 full	HNO ₃ to pH < 2; Cool to 4°C
Lead (SW 7421)	One 1 L P bottle	7/8 full	HNO ₃ to pH < 2; Cool to 4°C
Oil and Grease (SW 9071)	One 1 L G bottle (c)	7/8 full	H₂SO₄ to pH < 2; Cool to 4°C
Toxicity Characteristic (SW 1311) Leaching Procedure	One 1 L G bottle (c)	7/8 full	Cool to 4°C

NOTES:

⁽a) Hold times are presented in Section 1.8

⁽b) All containers to have Teflon®-lined lids except for vials for volatile organics which will have Teflon®-lined septa. Container codes: G = glass AG = amber glass P = polyethylene

⁽c) Samples chosen for MS/MSD and/or Laboratory Duplicate require double the number of containers indicated.

$$\frac{N}{1} \quad \frac{N}{2} \quad \frac{N}{3} \quad \frac{C}{4} \quad \frac{C}{5} \quad \frac{\#}{6} \quad \frac{\#}{7} \quad \frac{?}{8} \quad \frac{?}{9} \quad \frac{?}{10}$$

The "NNN" portion of the sample ID shall consist of the unique IRPIMS site code for the site. The "CC" portion of the sample ID shall consist of a two character location code, as follows:

MW - Monitoring well

SW - Surface water

SB - Soil boring in which no monitoring well was installed

SD - Sediment

SS - Surface soil

DC - Drill cuttings

The "##" portion of the sample ID shall consist of a number that is unique for each site and matrix, starting with "01." Characters 9 and 10 are optional for water samples and shall signify the upper depth of sample collection for soil and sediment samples. If characters 9 and/or 10 of the sample ID are used, character 8 shall be a dash. If characters 9 and 10 are not used, character 8 shall also not be used. Note that field duplicate samples will be given a false sample ID and date and time of sampling to hide their true identity from the laboratory.

For field QC samples, the following sample ID scheme shall be followed:

The "CC" portion of the sample ID shall consist of the appropriate two letter IRPIMS code for the type of QC sample, as follows:

AB - Ambient conditions blank

EB - Equipment blank

TB - Trip blank

Character 3 shall be a unique number for that QC sample type and for the day of collection, starting with "1." Character 4 shall be a dash. Characters 5 through 10 shall indicate the date of collection, where "DD" is the day, "MM" is the month (01=January, etc.), and "YY" is the last two digits of the year.

2.2.2.4 <u>Handling and Shipping</u> - The properly labeled and sealed sample containers will be placed in plastic "Ziploc" type bags and sealed. Bagged sample containers will be placed in the bottom of the cooler. Bagged sample containers will be arranged in the cooler so that they do not touch, and will be packed in double-bagged ice and additional packing material to prevent breakage. Samples should be packed so as to maintain a temperature of 4 degrees celsius during shipment.

Analytical Request forms, as shown on Figure 2-8, and chain-of-custody documents (see Section 2.2.3) will be sealed in air-tight plastic bags and taped to the inside of the cooler lid. The lid of the container will be taped shut and sealed with custody seals (see Section 2.2.3). Samples will be shipped directly to the laboratory by overnight courier on the day they are collected, if at all possible. The laboratory will be notified by phone of the sample shipment schedule. No samples will be held more than 24 hours on site, unless special arrangements are made with the laboratory.

2.2.3 Sample Custody

Documentation of sample custody from the time of collection will be maintained. Custody within the laboratory is described in Appendix B. Custody in the field is described in the following paragraphs. Chain-of-Custody records will be prepared for each shipping container. The custody record will be fully completed, in triplicate, by the field technician designated as responsible for

FIGURE 2-8 REQUEST FOR ANALYTICAL FORM

187155

	Project #: Manager:			
	SAMPLE ID:		_	
		ANALYSIS REC	QUEST	
Container	No.	Preservation	Parameter	Method
	COMMENT:		- <u>-</u>	

HF

PLEASE RETURN A COPY OF THIS FORM WITH THE DATA PACKAGE

sample shipment to the laboratory. The information specified on the chain-of-custody record will contain the same level of detail found in the site logbook, with the exception that on-site measurement data will not be recorded. The custody record will include the following information: name of person(s) collecting the samples, date samples were collected, the sample identification code (ID), type of sampling conducted (composite/grab), and number and type of containers used. Figure 2-9 presents an example of a Chain-of-Custody record.

Transfer of sample custody shall be indicated by the signatures of the sampler(s) and the person receiving the samples on the chain-of-custody record, with date and time noted. Prior to sealing the shipping container, the original and first copy of the chain-of-custody record shall be placed in a plastic bag that will be attached to the inside of the shipping container lid. The second copy remains with the sampler(s). If a courier service is to be used, the signature of the receiving person will be replaced by the shipper's airbill number. Receipt at the laboratory shall be indicated by signature of the laboratory's Sample Custodian, with the date and time recorded.

To allow a means of determining whether sample integrity was maintained, custody seals will be placed on the shipping containers. Custody seals will be pre-printed, adhesive-backed seals with security slots designed to break if disturbed. Sample shipping containers (coolers, cardboard boxes, etc., as appropriate) will be sealed in as many places as necessary to ensure security. Seals will be signed and dated before use. Upon receipt by the laboratory, the Sample Custodian will check and document whether the custody seals are intact and will also record the cooler temperature by measuring the temperature of the temperature vial. Any other sample integrity information will be recorded on the Chain of Custody and Cooler Receipt Forms.

SAMPLING INFORMATION STREET ADDRESS: CITY/STATE: CITY/STATE: O ON WHAT THE PROPESS: O ON WHAT THE PROPESS: O ON WHAT THE PROPESS: ATURE) SIGNATURE) SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNATURE SIGNAT			CHAIN	OF CUSTODY RECORD	TODY	RECOR	۵	4636	(0	1
S SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE SGNATURE S				SAMPLING INFORMATION	NAME OF FA STREET ADI CITY/STATE	ACILITY:		ZIP:		
S (SIGNATURE) S AMPLE STATION DESCRIPTION S SOLORE S SIGNATURE) S SIGNATURE S SIGNAT	PROJECT NAME		JOB NO.							
STATE DATE TIME RECEIVED BY: RELINQUISHED BY: RELINQUISHED BY: SIGNATURE S	SAMPLERS (SIGNATURE)			BNIATN	POAL SAN	///			//	\mathcal{I}
Colored Colo	SAMPLING DATE			OE CO	100000 NO.	(X)				7
FED BY: PATE / TIME RECEIVED BY: RELIMAUISHED BY: PATE / TIME RECEIVED BY: RELIMAUISHED BY: REL	GHAB COMP.	SAMPLE STATION	DESCRIPTION			&X,				$\overline{\mathcal{L}}$
FED PT. PATE / TIME RECEIVED BY: RELINQUISHED BY: RECEIVED BY: RELINQUISHED BY: RECEIVED BY:										
## FECHVED BY: DATE / TIME										
FED BY: PATE / TIME RECEIVED BY: FELINGUISHED BY: FECEIVED BY: FELINGUISHED BY: FECEIVED BY: FELINGUISHED BY: FELINGUISHE										
FELINOLISHED BY: PATE / TIME RECEIVED BY: FELINOLISHED BY: FELINOLISHED BY: GIGNATURE FECEIVED BY: GIGNATURE FELINOLISHED BY: GIGNATURE										
FIGURATURE PATE FRECEIVED BY: FRELINGUISHED BY: FACEIVED BY: FRELINGUISHED BY: FACEIVED BY: FAC					+					
FELINGUISHED BY: DATE / TIME RECEIVED BY: DATE / TIME RECEIVED BY: DATE / TIME RECEIVED BY: CSIGNATURE) CSIGNA										
ED BY: DATE / TIME RECEIVED BY: RELINQUISHED BY: CSIGNATURE) CSIGNATURE RECEIVED BY: CSIGNATURE RELINQUISHED BY: CSIGNATURE COVERY WELL - RW SOULSEDMIN - SO SUCSEDMIN - SO SUCSEDMIN - SO SULSEDMIN - SO SUCSEDMIN - SO SUCSE										
FECHINGUISHED BY: DATE / TIME RECEIVED BY: BELINGUISHED BY: DATE / TIME RECEIVED BY: FECHINGUISHED BY: FECHINGUISHED:										
IGANATURE) (SIGNATURE)	RELINQUISHED BY:	DATE / TIME	RECEIVED BY:		RELINQUISHED F	34:	DATE / TIME	RECEIVED BY:	1	4
FECHIVED BY: DATE / TIME RECEIVED BY: RELINQUISHED BY: DATE / TIME RECEIVED AT LABORATORY:	(SIGNATURE)	<u></u>	(SIGNA	TURE)	(SIGN	ATURE)		(SIGNATURE	.)	_
SIGNATURE) (SIGNATURE) (SIGNATURE) ION: ORIGINAL AND PINK COPIES ACCOMPANY SAMPLE SHIPMENT TO LABORATORY. * SOURCE RECOVERY WELL - RW PINK COPY RETAINED BY LABORATORY. * SOURCE SOIL SEDIMENT - SO SOIL SEDIMENT - SO SULSEDIMENT - SO SULSEDIM	RELINGUISHED BY:	DATE / TIME	RECEIVED BY:		RELINQUISHED F	3 Y :	DATE / TIME	RECEIVED AT LABORA	ATORY:	1.8
ION: ORIGINAL AND PINK COPIES ACCOMPANY SAMPLE SHIPMENT TO LABORATORY. PINK COPY RETAINED BY LABORATORY. YELLOW COPY RETAINED BY SAMPLERS. CODES: RCRA MONITORING WELL - RW SOIL/SEDIMENT - SO SLUGGE - SL NPDES DISCHARGE - ND	(SIGNATURE)		(SIGNA	(TURE)	(SIGN)	ATURE)	-	(SIGNATURE		37
	ä Ö	D PINK COPIES ACCOME ETAINED BY LABORATO	PANY SAMPLE SHIF	PMENT TO LABORATY Y RETAINED BY SAN	rory. Aplers.		sourc codes		WELL-MW	

2.2.4 OC Samples

Quality control (QC) samples are collected and analyzed for the purpose of assessing the quality of the sampling effort and the analytical data. QC samples include duplicates and replicates of field samples, equipment blanks, trip blanks, and ambient condition blanks. The type, description, preparation and collection of field QC samples are discussed in the following section.

- OC Samples Samples initiated and analyzed to identify and diagnose problems related to sampling analysis.
- Replicate A single sample divided into two equal parts for analysis in order to provide precision data. Replicates are applicable to soil and sediment samples.
- <u>Duplicate</u> Two samples collected independently at a sampling location during a single act of sampling in order to provide precision data. Duplicates are applicable to water samples.
- Equipment Blank Samples consisting of reagent water collected from a final rinse of sampling equipment after the decontamination procedure has been performed. The purpose of rinsate blanks is to determine whether the sampling equipment is causing cross contamination of samples.
- Trip Blank Containers of Type II reagent grade water that are kept with the field sample containers from the time they leave the laboratory until the time they are returned to the laboratory. The purpose of the trip blank is to determine whether samples are being contaminated during transit to and from the laboratory. Trip blanks pertain only to volatile organic analyses; therefore, the containers must contain no headspace. Only one trip blank is required per sampling day and shall satisfy trip blank requirements for all water

matrices for that day if the volatile samples are shipped in the same cooler. However, if more than one cooler contains volatile samples, then a trip blank will be placed in each cooler containing volatile samples.

Ambient Conditions Blank - Samples consisting of Type II reagent grade water that are prepared at the site. Ambient condition blanks are used to measure the influence of ambient volatile conditions at a particular set of sampling locations. Ambient condition blanks will not necessarily be collected every time volatile samples are collected. They will be collected when samples are collected downwind of possible volatile sources such as active runways or an adjacent highway.

2.2.5 Sample Analysis Summary

Tables summarizing the environmental samples and the proposed QC samples to be analyzed by parameter by site are as follows:

<u>Matrix</u>	<u>Table No.</u>
Subsurface Soil from borings	2-3
Ground Water - Monitoring Wells	2 - 4
Ground Water - Geochemical Survey	2-5
Surface Water	2-6
Sediments	2 - 7

2.3 FIELD MEASUREMENTS

2.3.1 Parameters for Field Characterization of Samples

Certain measurements must be performed as soon as possible after sampling in order to get accurate results. These parameters

include pH, temperature and specific conductivity. These parameters will be measured in the field when the samples are obtained during sampling. Other field parameters include safety measurements of air vapors and turbidity during well development.

2.3.2 Equipment Calibration

All equipment will be calibrated according to manufacturers instructions or a generally accepted practice. Calibration of all instruments will be recorded in the field book. The following subsections describe calibration procedures. Additional detail is provided in Appendix A.

- Measurement of pH will be done on site with a pH meter. The instrument will be field-calibrated with two buffer solutions at the beginning of each day's use. Accuracy of the measurement is maintained by selecting a standard buffer with a pH as close as possible to that of the sample (preferably within three pH units). The pH of the buffers used will be dependent upon the pH of the sample, but in each case will "bracket" the range of measurement. The stability of the calibration will be verified through the analysis of one standard periodically throughout the day as deemed necessary by the site manager, but at least once every 5 hours. Refer to Appendix A for calibration procedures.
- Temperature and specific conductivity will be measured with a portable meter. Calibration of the instrument is periodically performed at the factory as part of Law's internal QA program. The instrument probe will be rinsed with reagent water between each use and the calibration of the specific conductivity probe checked at the beginning and end of each day's use, using reagent grade water and two potassium chloride (KCl) solutions with known conductances that bracket the expected sample conductivities.

- The hand held portable organic vapor analyzer (OVA) with a photoionization detector (PID) used to screen the air vapors at the head of the augers and in the breathing zone will be calibrated daily with a 100 ppm level of isobutylene.
- Turbidity will be measured with a nephelometer (also known as a turbidimeter). The instrument will be calibrated with three standards of concentrations at 0.1, 0.2, and 0.5 NTUs at the beginning of each day's use. The stability of the calibration will be verified through the analysis of one standard periodically throughout the day as deemed necessary by the site manager, but at least once every 5 hours. A complete calibration procedure is described for the nephelometer in Appendix D.

2.3.3 Equipment Maintenance

Equipment used in the field is located in a central store room. The equipment must be "signed in" upon return and "signed out" when removed for use in the field. All equipment must be returned decontaminated and any malfunctions reported to the project manager. The project manager will incorporate any actions necessary for the repair or replacement of the equipment. Equipment maintenance logs are kept on file. Battery powered instruments will have their power supplies checked daily. Rechargeable instruments will be recharged daily.

2.3.4 Decontamination of Field Instruments

Decontamination of field instruments will depend upon the instrument. The probes for the pH, temperature and specific conductivity meters will be rinsed with reagent grade water before and after each use and at the end of each day. The measurement

vial for the turbidity meter will be rinsed out with deionized water before and after each use. No decontamination is required for the organic vapor analyzer.

2.4 FIELD OA/OC PROGRAM

2.4.1 Control Parameters

Control parameters of the field procedures consist of the same controls that govern analytical data. Control parameters consist of the following:

- · Collection of field and QC samples
- · Calibration of field equipment
- · Decontamination of field equipment
- · Development of wells
- · Purging a minimum of three well volumes before sampling
- · Record keeping

2.4.2 Control Limits and Corrective Actions

Appendix A presents a summary of field quality control procedures and acceptance criteria for field measurements. Corrective action requirements are presented in this summary.

2.5 RECORD KEEPING

Pertinent information concerning all aspects of sampling and field measurements will be recorded in hard-bound field notebooks, and all chain-of-custody documents completed in order to completely reconstruct the activities. The bound field notebook will have prenumbered pages, and entries will be made in indelible ink.

Environmental conditions will also be characterized. Each sampling site will be characterized by the following criteria.

- 1. Location of work
- 2. Weather
- 3. Rainfall
- 4. Temperature minimum and maximum
- 5. Wind direction
- 6. Ongoing activities that may influence or disrupt sampling efforts
- 7. Accessibility to the sampling locations, e.g., rough terrain, fallen trees, flooding, etc.

Each day the site manager will prepare a Daily Quality Control Report (DQCR) and send it to the AFCEE TC. The DQCR will include weather information at the time of sampling, samples collected, field instruments and calibrations and will reflect any problems that occurred in the field. A DQCR is shown in Figure 2-10.

2.6 SITE MANAGEMENT

The AFCEE Team Chief (TC) for the Carswell AFB RFI is Mr. Chris Hobbins. Mr. Hobbins' address and phone number are:

HQ AFCEE/ESB 8001 Inner Circle Dr., Suite 2 Brooks AFB, TX 78235-5000 (210) 536-5261

The Carswell AFB Point of Contact (POC) is Mr. Frank Grey. Mr. Grey's address and phone number are:

Air Force Base Conversion Agency Chief Environmental Engineer AF BCA/OL-H 1 Warehouse Rd. Carswell AFB, TX 76127-5000 (817) 782-6311

FIGURE 2-10 DAILY QUALITY CONTROL REPORT 187164

REPORT No	CONTRACT No		_ DATE	
LOCATION OF WORK				-
DESCRIPTION OF WORK _				
WEATHER	RAINFALL (INCHES)	TEMPERATURE _	MIN	MAX
WIND DIRECTION				
1. WORK PERFORMED				
3. PERSONNEL AND VISITO	ORS AT SITE			
	SITE MANAGER:		_	

2-55

HF

Carswell AFB personnel (Civil Engineering) will assist in base cooperation for the following:

- · Personnel identification and vehicle passes
- · Utility clearances and permits
- · Temporary field office
- A water supply for large quantities of potable water to be used in equipment cleaning, etc.
- Rights of easement and access to all Air Force and private property to perform all required field investigations

All field personnel will adhere to the procedures as described in the work plans. The site manager will supervise all field activities and continually update Mr. Hobbins and Law's Project Manager, John O'Brien, of the daily activities. Any major problems that occur during field activities will be brought to the attention of the TC for his input into the resolution of the problem. Emergency contacts for medical assistance are discussed in the Health and Safety Plan.

REFERENCES

- AFCEE, 1991. Handbook To Support The Installation Restoration

 Program (IRP) Statements of Work: Volume I Remedial

 Investigation/Feasibility Studies (RI/FS), United States Air

 Force, Air Force Center for Environmental Excellence [AFCEE]:

 Environmental Services Office, Environmental Restoration

 Division, Brooks Air Force Base, TX, May 1991 (Reprinted 22

 May 1992).
- AFCEE, 1993a. <u>Installation Restoration Program Information</u>

 <u>Management System (IRPIMS): Contractor Data Loading Tool</u>

 (CDLT) User's Manual Version 1.2, United States Air Force,

 Air Force Center for Environmental Excellence [AFCEE]:

 Environmental Services Office, Environmental Data Management

 Division, Brooks Air Force Base, TX, January 1993.
- AFCEE, 1993b. <u>Installation Restoration Program Information Management System (IRPIMS): Quality Control Tool (QCTOOL) User's Manual Version 1.2</u>, United States Air Force, Air Force Center for Environmental Excellence [AFCEE]: Environmental Services Office, Environmental Data Management Division, Brooks Air Force Base, TX, January 1993.
- ANSI 3.1, 1981, American National Standards Institute, Inc.,
 "American National Standard for Selection, Qualification, and
 Training of Personnel for Nuclear Power Plants."
- Bouwer, 1989. Bouwer, Herman, "The Bouwer and Rice Slug Test An Update," <u>Groundwater</u>, 1989, 27 (3 May-June), pp. 304-9.
- 42nd Strat Hosp/SGPB, 1988. Evaluation of Water From Demineralization Plant Ion Exchange, Water Analysis Letter to 42nd CES/DEEV, 12-12-88.

- USEPA, 1980. Interim Guidelines And Specifications For Preparing

 Quality Assurance Project Plans, (QAMS-005/80), United States

 Environmental Protection Agency [USEPA], Office of Monitoring

 Systems and Quality Assurance, Office of Research and

 Development, Washington, D.C., 29 December 1980.
- USEPA, 1987a. <u>Data Quality Objectives For Remedial Response</u>
 <u>Activities Development Process</u>, EPA/540/G-87/003, United
 States Environmental Protection Agency [USEPA], Office of
 Emergency and Remedial Response and Office of Waste Programs
 Enforcement, Office of Solid Waste and Emergency Response,
 Washington, D.C., March 1987.
- USEPA, 1987b. <u>Data Quality Objectives For Remedial Response</u>
 <u>Activities Example Scenario: RI/FS Activities at a Site with</u>
 <u>Contaminated Soils and Ground Water EPA/540/G-87/004</u>, United States Environmental Protection Agency [USEPA], Office of Emergency and Remedial Response and Office of Waste Programs Enforcement, Office of Solid Waste and Emergency Response, Washington, D.C., March 1987.
- USEPA, 1987c. Expanded Site Inspection (ESI) Transitional Guidance

 For FY 1988 (OSWER Directive 9345.1-02), United States

 Environmental Protection Agency [USEPA], Office of Emergency
 and Remedial Response, Washington, D.C., October 1987.
- USEPA, 1986. <u>Test Methods for Evaluating Solid Waste</u>, Third Edition, SW-846, Solid Waste and Emergency Response, Washington, D.C., November 1986.

TAB

APPENDIX A

APPENDIX A

QUALITY CONTROL REQUIREMENTS

AND

REPORTING LIMITS

APPENDIX A-1

QUALITY CONTROL REQUIREMENTS

FOR

CHEMICAL ANALYSIS

<u>Method</u>	<u>Parameter</u>
120.1	Specific conductance (field analysis)
150.1	pH (field analysis)
170.1	Temperature (field analysis)
+	Metals by GFAA/CVAA
6010	Metals by ICP
8240	Volatile Organic compounds by GC/MS
8020	Volatile Organic Compounds by GC
8270	Extractable Organics by GC/MS
8010	Purgeable Halocarbons
8015	Petroleum Hydrocarbons
9070	Oil and grease
418.1	Total Recoverable Petroleum Hydrocarbons
1311	Toxicity Characteristic Leaching Procedure

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

MOITO & SUITO SOCIO	NOTINE PALICIA	 Check system as per manufacturer's instructions. Check standard 	3. Replace instrument,	 Check standard. Check system as per manufacturer's instructions. 	3. Replace instrument.	1. Analyze 3rd aliquot of sample.	Z. Flag Data. Flag Data.
ACCEPTANCE CRITERIA		± 25 umhos/cm		± 25 umhos/cm		± 50 units	S detection limit
FREQUENCY		Prior to trip		At beginning and end of day		1 per day	1 per day
QUALITY CONTROL CHECK	INSTRUMENT OC:	Calibration		Calibration Stability	SAMPLE QC:	Duplicate	Rinsate
PARAMETER/ METHOD (a)	Specific Conductance	(EPA-600/4-79-020)					

⁽a) References:

[&]quot;SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

CORRECTIVE ACTION		 Check system as per manufacturer's instructions. Check standard. Replace instrument. 	1. Check standard. 2. Check system. 3. Recalibrate.		 Analyze 3rd aliquot of sample. Flag data. 	Flag data.
ACCEPTANCE CRITERIA		± 0.1 units	± 0.2 units		± 0.5 units	≤ detection limit
FREQUENCY		1 per day at two levels	1 per hour at two levels		1 per day	1 per day
QUALITY CONTROL CHECK	INSTRUMENT QC:	Calibration	Calibration Stability	SAMPLE QC:	Duplicate	Rinsate
PARAMETER/ METHOD (a)	pH PPA (Feel and Control	(EPA-600/4-79-020)				

(a) References:

"SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA RCRA Facility Investigation Carswell Air Force Base, Texas

PARAMETER/ METHOD (a)	QUALITY CONTROL CHECK	FREQUENCY	ACCEPTANCE CRITERIA	CORRECTIVE ACTION
Temperature	INSTRUMENT OC:			
1 / 0.1 (field analysis) (EPA-600/4-79-020)	Calibration	Prior to trip	± 2 °C	 Check against precision thermometer certified by MICT
				2. Replace instrument.
	SAMPLE QC:			
	Duplicate	1 per day	± 2 °C	 Analyze 3rd aliquot of sample. Flag data.

⁽a) References:

[&]quot;SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

187174

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

CORRECTIVE ACTION	Review lab QC data to determine if they are in control. If not in control, flag data. Use data to evaluate whether proper collection procedures were followed. If not, determine further corrective action.	Qualify data.		Terminate analysis, solve problem, recalibrate and reanalyze samples analyzed since last good CCV.	Terminate analysis, solve problem, recalibrate and reanalyze samples analyzed since last good CCB.	Redigest and reanalyze all samples less than 10X the MDL.	Perform post digest spike and qualify all associated data.
ACCEPTANCE CRITERIA	Above 10x detection limit, % RPD must be less than current control limits: Aqueous samples - RPD < 20% Non-aqueous samples - RPD < 35%	Less than reported detection limits		5 pt. calibration; Measured value within 10% of true value (20% for Hg)	Absolute value < MDL	Absolute value < MDL	See Appendix B
FREQUENCY	1 for every 10 field samples collected	1 for every 10 field samples collected		ICV - prior to analysis CCV - after every 10 samples and end of analytical batch	ICB - after initial calibration verification CCB - after every 10 samples and end of analytical batch	1 per batch of samples, minimum of 1 per 20 samples	1 per batch of samples, minimum of 1 per 20 samples
QUALITY CONTROL CHECK	<u>FIELD QC:</u> Duplicate	Rinsate (c)	LABORATORY QC:	Icv/ccv	ICB/CCB	Prep Blank	MS/MSD
PARAMETER/ METHOD (a)	Metals by GFAA and CVAA Arsenic (GFAA: 7060) Lead (GFAA: 7421) Selenium (GFAA: 7740) Mercury (CVAA: 7470/7471)						

⁽a) References:

[&]quot;SW-846" = Test Methods for Evaluating Solid Waste, SW-846, USEPA, November 1986
"EPA-600/4-79-020" = Methods for Chemical Analysis of Water and Wastes, EPA-600/4-79-020, USEPA, March 1979, Revised March 1983
"MHETL" = Maine Health and Environmental Testing Laboratory

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA RCRA Facility Investigation Carswell Air Force Base, Texas

Metals by ICP 6010 (SW-846)

PARAMETER/ METHOD (a)	QUALITY CONTROL CHECK	FREQUENCY	ACCEPTANCE CRITERIA	CORRECTIVE ACTION
ls by ICP	FIELD QC:			
(SW-846)	Duplicate	1 for every 10 field samples collected	No more than 4 target compounds, each with a concentration exceeding 3 times the method detection limit can be present.	Review lab QC data to determine if they are in control. If not in control, flag data. Use data to evaluate whether proper collection procedures were followed. If not, determine further corrective action.
	Rinsate (c)	1 for every 10 field samples collected	Above 10x detection limit, % RPD must be less than current control limits: Aqueous samples - RPD < 30% Non-aqueous samples - RPD < 40%	Qualify data.
	LABORATORY QC:			
	Initial (ICV) and Continuing (CCV) Calibration Verification	ICV - prior to sample analysis CCV - after every 10 samples and end of analytical batch	4 pt. calibration; Measured value within 10% of true value	Terminate analysis, solve problem, recalibrate and reanalyze samples analyzed since last good CCV.
	Initial (ICB) and Continuing (CCB) Calibration Blank	ICB - after initial calibration verification CCB - after every 10 samples and end of analytical batch	Absolute value ≤ Method Detection Limit (MDL)	Terminate analysis, solve problem, recalibrate and reanalyze samples analyzed since last good CCB.
	Prep Blank	1 per batch of samples, minimum 1 per 20 samples	Absolute value ≤ Method Detection Limit (MDL)	Redigest and reanalyze all samples greater than the MDL but less than 10x the blank concentration.
	MS/MSD	1 per batch of samples, minimum 1 per 20 samples	See Appendix B	Perform a post-digestion spike and qualify data appropriately.
	Laboratory Control Sample (LCS)	1 per batch of samples, minimum 1 per 20 samples	80 - 120% recovery: waters 75 - 125% recovery: sol/sed.	Rerun. If still out of control, solve problem and reanalyze batch.
	Interference Check	Beginning and end of run or per 8 hour shift	80-120% recovery	Terminate analysis, solve problem, recalibrate and reanalyze samples analyzed since last good ICS.

⁽a) References:

[&]quot;SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

187176

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

1	92	11.0	ی د			18
CORRECTIVE ACTION	Review lab QC data to determine if there is a laboratory problem. If not, and same compounds are found in field samples at similar concentrations, resample entire batch.	Review lab QC data to determine if there is a laboratory problem. If not, and same compounds are found in field samples at similar concentrations, resample entire batch.	Review lab QC data to determine if they are in control. If not in control, flag data. Use data to evaluate whether proper collection procedures were followed. If not, determine further corrective action.	Qualify data.	Tune instrument; repeat. Tune instrument; repeat. Recalibrate instrument.	Rerun continuing calibration. If still out of control, recalibrate instrument.
ACCEPTANCE CRITERIA	No more than 4 target compounds, each with a concentration exceeding 3 times the method detection limit can be present.	No more than 4 target compounds, each with a concentration exceeding 3 times the method detection limit can be present.	Above 10x detection limit, % RPD must be less than current control limits: Aqueous samples - RPD < 30% Non-aqueous samples - RPD < 40%	Less than reported detection limits	lon abundance criteria; see method See Method SW846 5 pt. calibration; SPCCs exceed 0.300; CCCs response factor deviates < 30% from average	SPCCs exceed 0.300 with the exception of bromoform which must exceed 0.250; CCCs response factor deviates < 25% from average of initial calibration
FREQUENCY	1 for each batch of samples shipped to laboratory	Collected when samples are collected downwind of possible volatile sources.	1 for every 10 field samples collected	1 for every 10 field samples collected	Every 12 hours of operation Every 12 hours of operation Prior to analysis and when continuing calibration fails criteria	Every 12 hours of operation
QUALITY CONTROL CHECK	FIELD OC: Trip Blank	Ambient Blank	Duplicate	Rinsate (c)	LABORATORY QC: Sensitivity Check with BFB Mass Calibration Initial Calibration	Continuing Calibration
PARAMETER/ METHOD (a)	Volatile Organic Compounds by GC/MS 8240 (SW-846)					

[&]quot;SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory (a) References:

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

CODDECTIVE ACTION	Qualify results <10 times that found in the blank.	Reanalyze sample; flag data. Reanalyze samples. If still out of control, qualify the data.
ACCEPTANCE CRITERIA		See Appendix B See Appendix B
FREQUENCY	1 for every 20 samples, or every Less than reported detection limit	Every sample 1 for every 20 samples
QUALITY CONTROL CHECK	Method Blank	Surrogate Recovery MS/MSD
PARAMETER/ METHOD (a)	Volade Organic Compounds Method Blank by GC/MS 8240 (SW-846) (Cont'd)	

⁽a) References;

[&]quot;SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes</u>, EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

TABLE A-1

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas **RCRA Facility Investigation**

PARAMETER/ METHOD (a)	QUALITY CONTROL CHECK	FREQUENCY	ACCEPTANCE CRITERIA	CORRECTIVE ACTION
Volatile Organic Compounds by GC 8020 (SW-846)	FIELD OC: Trip Blank	1 for each batch of samples shipped to laboratory	No more than 4 target compounds, each with a concentration exceeding 3 times the method detection limit can be present.	Review lab QC data to determine if there is a laboratory problem. If not, and same compounds are found in field samples at similar concentrations, resample entire batch.
	Ambient Blank	Collected when samples are collected downwind of possible volatile sources.	No more than 4 target compounds, each with a concentration exceeding 3 times the method detection limit can be present.	Review lab QC data to determine if there is a laboratory problem. If not, and same compounds are found in field samples at similar concentrations, resample entire batch.
	Duplicate	1 for every 10 field samples collected	Above 10x detection limit, % RPD must be less than current control limits: Aqueous samples - RPD < 30% Non-aqueous samples - RPD < 40%	Review lab QC data to determine if they are in control. If not in control, flag data. Use data to evaluate whether proper collection procedures were followed. If not, determine further corrective action.
	Rinsate (c)	1 for every 10 field samples collected	Less than reported detection limits	Qualify data.
	LABORATORY QC:			
	Initial Calibration	Prior to analysis and when continuing calibration fails criteria	5 pt. calibration: RSD ≤ 20% of RFs or correlation coefficient ≥ 0.995 (use avg. RF is RFD ≤ 20%, otherwise use calibration curve)	Recalibrate instrument.
	Continuing Calibration	Every 12 hours of operation	Measured value within 15% of true value	Rerun continuing calibration. If still out of control, recalibrate instrument.
	Method Blank	1 for every 20 samples, or every day	Less than reporting detection limit	Qualify results $<$ 10 times that found in the blank.
	Surrogate Recovery	Every sample	See Appendix B	Reanalyze sample; flag data.
	MS/MSD	1 for every 20 samples	See Appendix B	Reanalyze samples. If still out of control, qualify the data.

"SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory (a) References:

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas **RCRA Facility Investigation**

CORRECTIVE ACTION	Review lab QC data to determine if they are in control. If not in control, flag data. Use data to evaluate whether proper collection procedures were followed. If not, determine further corrective action.	Qualify data.	Tune instrument, repeat.	Tune instrument; repeat.	Recalibrate instrument.	Rerun continuing calibration; if still out of control, recalibrate instrument.	Reanalyze blank, then re-extract, qualify all results < 10 times that found in blank (for blanks contaminated with common laboratory chemicals) and qualify all results <5 times that found in blank (for blanks contaminated with compounds not commonly found in the laboratory).	Rerun sample. If still out of control, re-extract, reanalyze, qualify data.	Rerun sample. If still out of control, qualify data.
ACCEPTANCE CRITERIA	Above 10x detection limit, % RPD must be less than current control limits: Aqueous samples - RPD < 20% Non-aqueous samples - RPD < 35%	Less than reporting limit	lon abundance criteria; see method	See Method SW846	5 pt. calibration; SPCCs exceed 0.050; CCCs response factor deviates < 30% from average	SPCCs exceed 0.050; CCCs response factor deviates < 25% from average of initial calibration	Less than reporting detect limit	See Appendix B	See Appendix B
FREQUENCY	1 for every 10 field samples collected	1 for every 10 field samples collected	Every 12 hours of operation	Every 12 hours of operation	Before analysis and when continuing calibration fails criteria	Every 12 hours of operation	1 for every 20 samples	Every sample	1 for every 20 samples
QUALITY CONTROL CHECK	F <u>IELD QC:</u> Duplicate	Rinsate (c)	LABORATORY QC: Sensitivity Check with DFTPP	Mass Calibration	Initial Calibration	Continuing Calibration	Method Blank	Surrogate Recovery	MS/MSD
PARAMETER/ METHOD (a)	Extractable Organic Compounds by GC/MS 8270 (SW-846)								

"SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory (a) References:

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

	CORRECTIVE ACTION	Review lab QC data to determine if they are in control, If not in control, flag data. Use data to evaluate whether proper collection procedures ware	followed. Qualify data.	Qualify data.		Recalibrate instrument.	Rerun continuing calibration: if still out of control	recalibrate instrument. Step 1: Reanalyze Step 2: If second blank exceeds criteria, clean the analytical system. Step 3: Document the correction of the c	qualify all associated data. Rerun samples. If still out of control, qualify data.	Rerun. If still out of control, solve problem and reanalyze batch.
ACCEPTANCE CRITERIA	ליינים המיינים	Water samples - RPD < 20% Soil samples - RPD < 30%	Less than reporting detection limits	Less than reported detection limit		5 pt. calibration %RSD must be < 20% for all analytes	Response factor must be < 15%	rrom average of initial calibration Compounds must be below respective detection limits	See Table 1-32 for current control limits	Aqueous MS/MSD criterion apply
FREQUENCY		1 for every 10 field samples collected	1 for every 10 field samples collected	1 for each batch of samples shipped to laboratory		Prior to analysis and when continuing calibration fails criteria	Daily	Daily	1 per every 20 samples	1 per every 20 samples
QUALITY CONTROL CHECK	FIELD QC:	Duplicate	Rinsate	Trip Blank (applies to volatile fraction only)	LABORATORY QC:	Initial Calibration	Continuing Calibration	Method Blank	MS/MSD	rcs
METHOD (a)	SW846 8015M	lotal Petroleum Hydrocarbons (TPH)								

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

		CORRECTIVE ACTION	Review (ahe) OC docs to 1	Review label QC data to determine if they are in control. If not, qualify data. Use data to evaluate proper collection procedures were followed. If not, determine further corrective action.				reanalyze all samples greater than MDL but less than 10x blank concentration.	Rerun. If still out of control, reanalyze. Qualify data.
	ACCEPTANCE CRITERIA	WIND TOWN	% RPD must be ≤ 40%		Less than reported method detection	limits	Value must be less than sented	detection limit	See Table 1-32 for current control limits
	FREQUENCY		1 for every 10 field samples		1 for every 10 field samples			minimum 1 per 20 samples	i per matrix type, minimum 1 per 20 samples
QUALITY CONTROL	CHECK	FIELD QC:	Duplicate		Rinsates	LABORATORY OC:	Blank	MS/MSD	
PARAMETER/ METHOD (a)		SW 1311 TCLP							

(a) References:

"SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility Investigation

CORRECTIVE ACTION		Review lab QC data to determine if there is a laboratory problem. If not, review field decontamination procedures to either validate or invalidate data. If invalid, determine further corrective action.	Review lab QC data to determine if they are in control. If not in control, flag data. Use data to evaluate whether proper collection procedures were followed. If not, determine further corrective action.		Rerun calibration standards.	Document and report to client.	Rerun spike; if still out of control, reanalyze spike.	Reprep duplicates and reanalyze.
ACCEPTANCE CRITERIA		Less than reporting detection limit	Above 10x detection limit, % RPD must be less than current control limits: Aqueous samples - RPD < 30% Non-aqueous samples - RPD < 40%		0.995 correlation with a 3 point calibration	Less than reporting detection limit	Within current control limits	Above 10x detection limit; see Table 13-1 for %RSD
FREQUENCY		1 for every 10 field samples	1 for every 10 field samples		Each batch	1 for every 20 samples or extraction batch	1 for every 20 samples	1 for every 20 samples
QUALITY CONTROL CHECK	FIELD QC:	Equipment Blank	Duplicate	LABORATORY QC	Calibration Curve	Method Blank	Spiked Sample	Duplicate Spike
PARAMETER/ METHOD (a)	SW 9071/EPA 418.1 Petroleum Hydrocarbons							

⁽a) References:

[&]quot;SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas **RCRA Facility Investigation**

CORRECTIVE ACTION		Review lab QC data to determine if there is a laboratory problem. If not, and same compounds are found in field samples at similar concentrations, resample entire batch.	Review lab QC data to determine if they are in control. If not in control, flag data. Use data to evaluate whether proper collection procedures were followed.	Qualify data.		Recalibrate instrument.	Rerun continuing calibration. If still out of control, recalibrate instrument.	Step 1: Reanalyze. Step 2: If second blank exceeds criteria, clean the analytical system. Step 3: Document the corrective action taken and qualify all associated data.	Step 1: Reanalyze. Step 2: If recovery still outside control limits, qualify the data.	Rerun samples. If still out of control, qualify data.
ACCEPTANCE CRITERIA		No more than 4 target compounds each with a concentration exceeding 3 times the method detection limit can be present.	Water Samples - RPD < 20% Soil Samples - RPD < 30%	Less than reported detection limits		5 pt. calibration; % RSD must be < 20% for all analytes	Response factors must be < 15% from average of initial calibration	Compounds must be below respective detection limits	See Appendix D for current control limits	See Appendix D for current control limits
FREQUENCY		1 for each batch of samples shipped to laboratory	1 for every 10 field samples collected	1 for every 10 field samples collected		Prior to analysis and when continuing calibration fails criteria	Daily	Daily	All blanks, standards, QC samples, field samples	1 per every 20 samples See Ap
QUALITY CONTROL CHECK	FIELD, QC:	Trip Blank	Duplicate	Rinsate (c)	LABORATORY QC:	Initial Calibration	Continuing Calibration	Method Blank	Surrogate Spike	MS/MSD 1 per every
PARAMETER/ METHOD (a)	SW846 8010 Halogenated Volatile Organics									WS

⁽a) References:

[&]quot;SW-846" = <u>Test Methods for Evaluating Solid Waste</u>, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory

SUMMARY OF QUALITY CONTROL REQUIREMENTS AND ACCEPTANCE CRITERIA Carswell Air Force Base, Texas RCRA Facility investigation

CORRECTIVE ACTION		Review lab QC data to determine if they are in control. If not, qualify data. Use data to evaluate proper collection procedures were followed. If not, determine further corrective action.	Qualify data.		Check balance maintenance, qualify data.	Reanalyze all samples greater than MDL but less than $10 \times \text{blank}$ concentration.	Qualify data.
ACCEPTANCE CRITERIA		% RPD <u><</u> 35%	Less than reported detection limits		All measures must be accurate.	Less than reported detection limits	RPD ≤ 20%
FREQUENCY		1 for every 10 field samples collected	1 for every 10 field samples collected		According to the instrument service manual	1 per batch of 20 samples	1 per batch of 20 samples
QUALITY CONTROL CHECK	FIELD QC:	Duplicate	Rinsate (c)	LABORATORY QC:	Calibration of the instrument	Method Biank	Duplicates
PARAMETER/ METHOD (a)	SW9070						

[&]quot;SW-846" = Test Methods for Evaluating Solid Waste, SW-846, USEPA, November 1986 "EPA-600/4-79-020" = <u>Methods for Chemical Analysis of Water and Wastes,</u> EPA-600/4-79-020, USEPA, March 1979, Revised March 1983 "MHETL" = Maine Health and Environmental Testing Laboratory (a) References:

Table A-2 PROJECT REQUIRED REPORTING LIMITS - SOIL/SEDIMENT LAW ENVIRONMENTAL, INC. - NATIONAL LABORATORIES - PENSACOLA (LENL-P) June 1993

			June 1993			CONTROL	LIMITS
*****				RTING			RELATIVE %
ANALYTICAL				MIT	SPIKE	RECOVERY	DIFFERENCE
METHOD_	COMPOUND	UNITS	<u>IDL</u>	MQL	CONCENTRATION	<u> </u>	%
6010 (SW846)	Aluminum	mg/kg	10	50	*	75-125 **	20
	Antimony	mg/kg	5.6	15	50	78-117	18
	Arsenic	mg/kg	4.2	30	200	75-123	19
	Barium	mg/kg	3	10	200	85-121	20
	Beryllium	mg/kg	0.2	1	5	80-116	20
	Cadmium	mg/kg	0.4	0.5	5	87-123	20
	Calcium	mg/kg	10.2	100	*	75-125 **	20
	Chromium	mg/kg	0.8	5	20	82-124	18
	Cobalt	mg/kg	1.3	5	50	88-124	17
	Copper	mg/kg	0.8	5	25	89-125	20
	Iron	mg/kg	5	5	*	75-125 **	20
	Lead	mg/kg	3.7	5	50	76-125	18
	Magnesium	mg/kg	11.7	100	*	75-125 **	20
	Manganese	mg/kg	0.6	2	50	88-124	20
	Molybdenum	mg/kg	3.6	10	50	75-125 **	20
	Nickel	mg/kg	2.5	15	50	85-125	19
	Potassium	mg/kg	13.4	100	*	75-125 **	20
	Selenium	mg/kg	7.9	50	200	76-124	20
	Silver	mg/kg	0.8	5	5	80-125	18
	Sodium	mg/kg	25.5	100	*	75-125 **	20
	Thallium	mg/kg	7	7	200	76-121	20
	Vanadium	mg/kg	1 1	10	50	81-123	19
	Zinc	mg/kg	0.7	2	50	80-122	20
				_ <u>-</u>			
			PQL				
8240 (SW846)	Acetone	mg/kg	0.001	0.01	0.05	D-208	40
	Benzene	mg/kg	0.001	0.001	0.05	32 – 171	40
	Bromodichloromethane	mg/kg	0.001	0.001	0.05	16-198	40
	Bromoform	mg/kg	0.005	0.005	0.05	30-186	40
	Bromomethane	mg/kg	0.001	0.001	0.05	9-213	40
	2-Butanone (MEK)	mg/kg	0.01	0.01	0.05	D-213	40
	Carbon disulfide	mg/kg	0.005	0.005	0.05	D-222	40
	Carbon tetrachloride	mg/kg	0.005	0.005	0.05	34-161	40
	Chlorobenzene	mg/kg	0.005	0.005	0.05	20-186	40
	Chloroethane	mg/kg	0.005	0.005	0.05	19-208	40
	2 - Chloroethyl vinyl ether	mg/kg	0.01	0.01	0.05	D-278	40
	Chloroform	mg/kg	0.005	0.005	0.05	27-152	40
	Chloromethane	mg/kg	0.001	0.001	0.05	18-201	40
	Dibromochloromethane	mg/kg	0.005	0.005	0.05	30-179	40
	1,1-Dichloroethane	mg/kg	0.005	0.005	0.05	37-162	40
	1,2-Dichloroethane	mg/kg	0.005	0.005	0.05	25-192	40
	1,1 - Dichloroethene	mg/kg	0.005	0.005	0.05	7-229	40
	trans-1,2-Dichloroethene	mg/kg	0.005	0.005	0.05	51-143	40
	1,2-Dichloropropane	mg/kg	0.001	0.001	0.05	2-225	40
	cis-1,3-Dichloropropene	mg/kg	0.001	0.001	0.05	8-233	40
	trans-1,3-Dichloropropene	mg/kg	0.005	0.005	0.05	1-206	40
	Ethylbenzene	mg/kg	0.005	0.005	0.05	24 – 192	40
	2-Hexanone	mg/kg	0.005	0.00	0.05	D-229	40
	Methylene chloride	mg/kg	0.005	0.005	0.05	D-192	40
	i Metrivierie Chioride			0.000	0.00	D 102	
			0.01	0.01	0.05	D-226	40
	4-Methyl-2-pentanone (MIBK)	mg/kg	0.01	0.01	0.05 0.05	D-226	40
			0.01 0.005 0.005	0.01 0.005 0.005	0.05 0.05 0.05	D-226 6-196 24-179	40 40 40

NA = Not Applicable

NS = Not Spiked

NE = Not Established at this time

MQL = Maximum Quantitation Limit

^{**} Project – Required Reporting Limits for methodologies performed by TEG are listed in the TEG QA Manual in Appendix B – 2.

** = Precision and accuracy limits are based on method limits

IDL = Investment Detection Limit

Table A-2 PROJECT REQUIRED REPORTING LIMITS - SOIL/SEDIMENT LAW ENVIRONMENTAL, INC. - NATIONAL LABORATORIES - PENSACOLA (LENL-P) June 1993

			June 1993			CONTROL	LIMITS
ANALYTICAL				ORTING MIT	SPIKE	RECOVERY	RELATIVE % DIFFERENCE
METHOD	COMPOUND	UNITS	IDL	MQL	CONCENTRATION	%	%
8240 (SW846)	Toluene	mg/kg	0.005	0.005	0.05	18-177	40
	1,1,1-Trichloroethane	mg/kg	0.005	0.005	0.05	31-174	40
	1,1,2-Trichloroethane	mg/kg	0.005	0.005	0.05	29-171	40
	Trichloroethene	mg/kg	0.005	0.005	0.05	37-171	40
	Vinyl acetate	mg/kg	0.01	0.01	0.05	5-239	40
	Vinyl chloride	mg/kg	0.002	0.002	0.05	6-215	40
	Xvienes (total)	mg/kg	0.005	0.005	0.05	7-211	40
	1,2-Dichloroethane-d4 (surrogate)	mg/kg	NA	NA	0.05	72-117	NA
	Toluene - d8 (surrogate)	mg/kg	NA.	NA	0.05	87-116	NA
	4 - Bromofluorobenzene (surrogate)	mg/kg	NA	NA	0.05	75-107	NA
8020 (SW846)	Benzene	mg/kg	0.0005	0.0005	0.01	23-171	40
	Toluene	mg/kg	0.0005	0.0005	0.01	61-142	40
	Ethylbenzene	mg/kg	0.0005	0.0005	0.01	15-178	40
	Xylenes	mg/kg	0.001	0.001	0.03	50-156	40
	Fluorobenzene (surrogate)		NA	NA	0.03	77 – 125	NA NA
		mg/kg	NA NA	ł .			NA NA
6040 (C\4046\	4-Chlorotoluene (surrogate)	mg/kg	0.1	NA O.F.	0.04	79-124 77-123	20
6010 (SW846)		mg/L	1	0.5	2000		
	Antimony	mg/L	0.056	0.5	500	78-121	15
	Arsenic	mg/L	0.042	0.3	2000	78-122	17
	Barium	mg/L	0.03	0.1	2000	75-124	20
	Beryllium	mg/L	0.002	0.01	50	76-124	16
	Cadmium	mg/L	0.004	0.005	50	80-122	18
	Calcium	mg/L	0.102	1	NS	75-125 **	20
	Chromium	mg/L	0.008	0.05	200	76-121	15
	Cobalt	mg/L	0.013	0.05	500	75-123	14
	Copper	mg/L	0.008	0.05	250	76-124	18
	Iron	mg/L	0.05	0.05	1000	81 – 121	20
	Lead	mg/L	0.037	0.2	500	77-122	17
	Magnesium	mg/L	0.117	1	NS	75-125 **	20
	Manganese	mg/L	0.006	0.02	500	78-125	17
	Molybdenum	mg/L	0.036	0.1	500	75-123	16
	Nickel	mg/L	0.025	0.15	500	75-123	18
	Potassium	mg/L	0.134	5	NS	75-125 **	20
	Selenium	mg/L	0.079	1	2000	76-124	17
	Silver	mg/L	0.008	0.05	50	75-123	17
	Sodium	mg/L	0.255	1	NS	75-125 **	20
	Theilium	mg/L	0.072	0.4	2000	75-123	17
	Vanadium	mg/L	0.01	0.1	500	77-125	18
	Zinc	mg/L	0.007	0.02	500	78-125	16
7060 (SW846)	Arsenic	mg/L	0.002	0.005	40	75-118	18
7421 (SW846)	Lead	mg/L	0.002	0.005	20	75-123	16
7740 (SW846)	Selenium	mg/L	0.001	0.005	10	79-125	18
7471 (SW846)	Mercury	mg/L	0.0002	0.003	1	78-123	19
74/1 (011040)	IVIET CUT Y	mg/c	0.0002	0.001	<u> </u>	70-123	19
	 	PQL	+	 	+		+
8270 (SW846)	Benzoic acid	μg/L	36	50	50	D-500	40
(acids)	4-Chloro-3-methylphenol	μg/L	0.8	10	50	25-144	40
	2-Chlorophenol	μg/L	1	10	50	23-128	40
	2,4-Dichlorophenol	μg/L	0.9	10	50	40-124	40
	2,4 – Dimethylphenol	μg/L	1	10	50	38-126	40
	4,6-Dinitro-2-methylphenol	μg/L μg/L	0.6	50	50	D-166	40
	2,4-Dinitrophenol	μg/L μg/L	3.3	50	50	D-190	40
	2-Methylphenol	, -	0.8	10	50	22 – 147	40
		μg/L		1)		
	4-Methylphenol	μg/L	0.9	10	50	22-147	40

NA = Not Applicable
NS = Not Spiked

NE = Not Established at this time

IDL = Investment Detection Limit
MQL = Maximum Quantitation Limit

^{*} Project – Required Reporting Limits for methodologies performed by TEG are listed in the TEG QA Manual in Appendix B-2.

^{** =} Precision and accuracy limits are based on method limits

Table A-2 PROJECT REQUIRED REPORTING LIMITS - SOIL/SEDIMENT LAW ENVIRONMENTAL, INC. - NATIONAL LABORATORIES - PENSACOLA (LENL-P)

			lune 1993			CONTROL	LIMITS
		REPORTING		}	OOM	RELATIVE %	
ANALYTICAL		1	LIN		SPIKE	RECOVERY	DIFFERENCE
METHOD	COMPOUND	UNITS	IDL	MQL	CONCENTRATION	%	%
3270 (SW846)	2-Nitrophenol	μg/L	0.9	10	50	10-181	40
acids)	4 - Nitrophenol	μg/L	2	50	50	D-128	40
,,	Pentachlorophenol	μg/L	0.6	30	50	10-171	40
	Phenol	μg/L	0.8	10	50	11-83	40
	2,4,5-Trichlorophenol	μg/L	1	50	50	D-250	40
	2,4,6 - Trichlorophenol	μg/L	0.7	10	50	41-142	40
	2-Fluorophenol (surrogate)	μg/L	NA	NA.	100	25-99	NA
	Phenol-d6 (surrogate)		NA.	NA.	100	17-93	NA NA
		μg/L	NA I	NA NA	100	16-118	NA NA
2070 (01110.40)	2,4,6-Tribromophenol (surrogate)	μg/L	0.4	10	50	56-135	40
8270 (SW846)	Acenaphthene	μg/L					
(base/neutrai)	Acenaphthylene	μg/L	0.6	10	50	40-137	40
	Anthracene	μg/L	0.6	10	50	40-131	40
	Benzo(a)anthracene	μ g/ L	0.4	10	50	36-133	40
	Benzo(b)fluoranthene	μg/L	0.8	10	50	31 – 148	40
	Benzo(k)fluoranthene	μ g /L	0.6	10	50	17-14 9	40
	Benzo(ghi) perylene	μ g /L	0.6	10	50	8-187	40
	Benzo(a) pyrene	μg/L	0.3	10	50	18-148	40
	Benzyl alcohol	μg/L	1.7	10	50	D-250	40
	bis(2-Chloroethoxy)methane	μg/L	0.5	10	50	34-163	40
	bis(2-Chloroethyl)ether	μg/L	0.9	10	50	19-147	40
	bis (2 - Chloroisopropyl) ether	μg/L	0.6	10	50	42-154	40
	bis(2-Ethylhexyl) phthalate	μg/L	1.3	10	50	15-149	40
	4-Bromophenyi phenyi ether	μg/L	0.6	10	50	55-116	40
	Butylbenzylphthalate	μg/L	0.5	10	50	52-124	40
	4-Chloroaniline	μg/L	4.8	10	50	D-250	40
			0.4	10	50	63-112	40
	2-Chloronaphthalene	μg/L	0.4	10	50	34-144	40
	4-Chlorophenyl phenyl ether	μg/L	•		1		40
	Chrysene	μg/L	0.5	10	50	23153	,
	Dibenzo(a,h)anthracene	μg/L	0.6	10	50	6-194	40
	Dibenzofuran	μg/L	0.4	10	50	D-250	40
	Di-n-butylphthalate	μg/L	1	10	50	16-116	40
	1,2-Dichlorobenzene	μg/L	0.7	10	50	34-125	40
	1,3-Dichlorobenzene	μg/L	0.6	10	50	5-160	40
	1,4-Dichlorobenzene	μg/L	0.7	10	50	21-112	40
	3,3'-Dichlorobenzidine	μg/L	2	20	50	14-213	40
	Diethylphthalate	μg/L	0.5	10	50	14-97	40
	Dimethylphthalate	μg/L	0.3	10	50	D-112	40
	2.4 - Dinitrotoluene	μg/L	0.8	10	50	35-124	40
	2.6Dinitrotoluene	μg/L	0.8	10	50	55-143	40
	Di-n-octylphthalate	μg/L	0.7	10	50	3-143	40
	Fluoranthene	μg/L	0.6	10	50	30-130	40
	Fluorene	1 !	0.5	10	50	59-119	40
	(μg/L	0.5	10	50	6-144	40
	Hexachlorobenzene	μg/L					
	Hexachlorobutadiene	μg/L	0.7	10	50	24-111	40
	Hexachlorocyclopentadiene	μg/L	1.8	10	50	7-86	40
	Hexachloroethane	μg/L	0.9	10	50	41 100	40
	Indeno[1,2,3-cd]pyrene	μg/L	1.6	10	50	2-140	40
	Isophorone	μg/L	0.5	10	50	26-196	40
	2-Methylnaphthalene	μg/L	0.5	10	50	D-250	40
	Naphthalene	μ g/ L	0.5	10	50	32-123	40
	2-Nitroaniline	μg/L	0.7	50	50	D-250	40
	3-Nitroaniline	μg/L	1	50	50	D-500	40
	4-Nitroaniline	μg/L	4	50	50	D-500	40
	Nitrobenzene	μg/L	0.9	10	50	42-160	40

NA = Not Applicable NS = Not Spiked

NE = Not Established at this time

IDL = Investment Detection Limit

MQL = Maximum Quantitation Limit

^{*} Project - Required Reporting Limits for methodologies performed by TEG are listed in the TEG QA Manual in Appendix B-2.

^{** =} Precision and accuracy limits are based on method limits

Table A-2 PROJECT REQUIRED REPORTING LIMITS - SOIL/SEDIMENT LAW ENVIRONMENTAL, INC. - NATIONAL LABORATORIES - PENSACOLA (LENL-P)

						CONTROL	
				RTING			RELATIVE %
ANALYTICAL				AIT	SPIKE	RECOVERY	DIFFERENCE
METHOD	COMPOUND	UNITS	IDL	MQL	CONCENTRATION	<u>%</u>	%
3270 (SW846)	n-Nitrosodiphenylamine	μ g/ L	1	10	50	13-143	40
base/neutral)	n-Nitrosodipropylamine	μ g/ L	1	10	50	9-171	40
	Phenanthrene	μg/L	0.4	10	50	55-113	40
	Pyrene	μ g /L	0.7	10	50	60-113	40
	1,2,4-Trichlorobenzene	μg/L	0.5	10	50	46-140	40
	Nitrobenzene – d5 (surrogate)	μg/L	NA	NA	50	39-104	NA
	2-Fluorobiphenyl (surrogate)	μ g/ L	NA	NA	50	46-110	NA NA
	Terphenyl-d14 (surrogate)	μg/L	NA _	NA NA	50	38-137	NA NA
			PQL				
8240 (SW846)	Acetone	μg/L	7.5	10	50	32-199	40
02 10 (0 110 10)	Benzene	μg/L	0.4	5	50	46-147	40
	Bromodichloromethane	μg/L	0.3	5	50	49-150	40
	Bromotorm	μg/L	0.2	5	50	66-163	40
	Bromomethane		0.2	10	50	28-156	40
	4	μg/L	2.8	10	50	29-156	40
	2-Butanone (MEK)	μ g /L			50		40
	Carbon disulfide	μ g/ L	0.7	5		21-184	
	Carbon tetrachloride	μg/L	0.4	5	50	78-132	40
	Chlorobenzene	μg/L	0.2	5	50	49-152	40
	Chloroethane	μg/L	0.3	10	50	35-177	40
	2-Chloroethyl vinyl ether	μg/L	8	10	50	9-244	40
	Chloroform	μg/L	0.3	5	50	56-126	40
	Chloromethane	μg/L	0.6	10	50	32-162	40
	Dibromochloromethane	μg/L	0.3	5	50	60-143	40
	1,1 - Dichloroethane	μg/L	0.3	5	50	68-139	40
	1.2-Dichloroethane	μg/L	0.4	5	50	29-153	40
	1.1 - Dichloroethene	μg/L	0.3	5	50	32-192	40
	trans-1,2-Dichloroethene	μg/L	0.4	5	50	81-119	40
	1,2-Dichloropropane	μg/L	0.6	5	50	12-192	40
	cis-1,3-Dichloropropene	μg/L	0.6	5	50	19-194	40
	trans-1,3-Dichloropropene	μg/L	0.2	5	50	19~177	40
	Ethylbenzene	μg/L	0.2	5	50	47-155	40
	2 – Hexanone		2.1	10	50	17-191	40
		μg/L	4.5	5	50	8-173	40
	Methylene chloride	μg/L					
	4-Methyl-2-pentanone (MIBK)	μg/L	7	10	50	19-186	40
	Styrene	μg/L	0.5	5	50	27-148	40
	1,1,2,2-Tetrachloroethane	μg/L	0.7	5	50	49-140	40
	Tetrachloroethene	μg/L	0.2	5	50	71 – 142	40
	Toluene	μg/L	0.3	5	50	60-144	40
	1,1,1—Trichloroethane	μg/L	0.4	5	50	5 6-150	40
	1,1,2-Trichloroethane	μ g /L	0.7	5	50	61 – 140	40
	Trichloroethene	μg/L	0.2	5	50	77 – 139	40
	Vinyl acetate	μg/L	0.7	10	50	26-200	40
	Vinyl chloride	μg/L	0.6	10	50	26-188	40
	Xylenes (total)	μg/L	0.7	5	150	14-181	40
	1,2-Dichloroethane-d4 (surrogate)	μg/L	NA	NA	50	77-113	NA
	Toluene – d8 (surrogate)	μg/L	NA.	NA.	50	90-110	NA.
	4-Bromofluorobenzene (surrogate)	μg/L	NA NA	NA NA	50	87-114	NA NA
8020 (SW846)	Benzene	μg/L	0.5	1	NE	38-155	40
5520 (511640)	Toluene		0.5	2	NE NE	76-125	40
	Ethylbenzene	μg/L		I	NE NE	76-125 32-159	40
		μg/L	0.5	4			
	Xylenes	μg/L	1	2	NE	67-138	40
	Fluorobenzene (surrogate)	μg/L	NA	NA NA	NE	NE	NA NA
	4-Chlorotoluene (surrogate) Total Recoverable Petroleum Hydrocarbo	μg/L	0.001	0.001	NE 41	NE 80-120	NA

NA = Not Applicable
NS = Not Spiked

NE = Not Established at this time

MQL = Maximum Quantitation Limit

Project – Required Reporting Limits for methodologies performed by TEG are listed in the TEG QA Manual in Appendix B – 2.

 *** = Precision and accuracy limits are based on method limits

IDL = Investment Detection Limit

TAB

APPENDIX B

APPENDIX B

QUALITY ASSURANCE PROGRAMS

FOR

CHEMICAL ANALYSIS LABORATORIES

- B-1 Law Environmental, Inc. National Laboratories Branch (LENL), Pensacola, Florida
- B-2 Transglobal Environmental Geochemistry, Texas

APPENDIX B-1

QUALITY ASSURANCE PROGRAMS

Law Environmental, Inc.
National Laboratories Branch (LENL)
Pensacola, Florida



Law Environmental, Inc.

187193

INTEROFFICE MEMORANDUM

TO:

Chemistry Department

FROM:

Jerry S. Preston 25th

DATE:

September 28, 1992

SUBJECT:

Certifications held by LENL-P Laboratory

Attached please find a listing of the certifications held by the LENL-Pensacola laboratory. This list will be useful during proposal and planning activities.

CERTIFICATION BY STATE

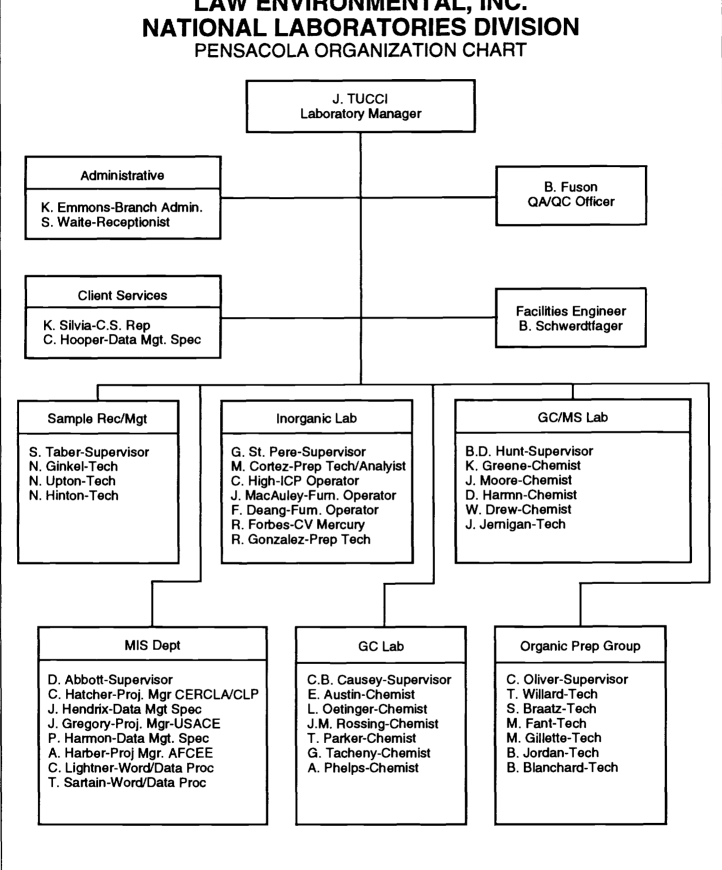
<u>STATE</u>	<u>PARAMETERS</u>	CERT. NO.	EXP. DATE
Alabama	Drinking Water	40600	6/30/93
Florida	Drinking Water	HRS#81351	6/30/93
Florida	Environmental Water	HRS#E81234	6/30/93
Florida	Water	HRS#81351	6/30/93
Kansas	Drinking Water and/or Pollution Control Sa	E-184 mples	7/31/93
Kansas	Solid or Hazardous Waste Samples	E-1161	7/31/93
Louisiana	Drinking Water	92-16	12/31/92
North Carolina	Wastewater	287	12/31/92
South Carolina	Wastewater	96008	Exp/Pending

<u>STATE</u>	<u>PARAMETERS</u>	CERT. NO.	EXP. DATE
Tennessee	BTX and TPH		7/01/93
Virginia	Water	00170	6/30/93
COE	Soil and Groundwater		2/20/94
USAF	Soil and Groundwater		9/25/93
U.S. Navy	Soil and Groundwater	Pending	
U.S. EPA	SAS Contracts	68-D9-0135	9/19/94

cc: Becky Pridgeon

/dsl

LAW ENVIRONMENTAL, INC.



LENL KEY PERSONNEL AND RESPONSIBILITIES

Position/Key Personnel	Responsibilities
Laboratory Manager James M. G. Tucci	Implement methodologies and procedures consistent with the generation of legally defensible data;
	 Provide overall management and operation of the lab;
	 Provide a safe working environment for employees;
	 Provide resolutions to items requiring corrective actions;
	 Provide training programs for employees;
	 Interact with QA/QC Coordinator to resolve analytical, methodological and QA/QC problems;
	- Obtain laboratory accreditations;
	 Schedule work in a manner consistent with personnel and instrumentation;
	 Implement procedures consistent with and building upon our philosophy of honesty, trust and ethics; and
	 Implement actions required to establish our lab as a premier laboratory.

3517-0111.07 1 of 5

Position/Key Personnel	Responsibilities
QA/QC Coordinator	Implement the branch OA program:
Burnie D. Fuson	- Implement the branch QA program;
	 Develop the QA/QC manual;
	- Establish and maintain safety
	standards and operating procedures;
	 Coordinate and audit the technical review of deliverables;
	- Issue recommendations and corrective actions required for any aspect of laboratory operations inconsistent with established policies and procedures;
	 Monitor and identify out-of-control or potentially out-of-control situations to Operations Manager, Supervisors, and Branch Manager;
	 Provide guidance for the chemistry QA/QC program development;
	 Interact with external QA personnel concerning the lab's certifications and QA policies/procedures and coordinating QA compliance as required;
	 Keep abreast of new techniques and programs for QA and safety and inform Branch Manager and Operations Manager;
	 Coordinate the development of project QA plans as required;
	 Provide historical QA reports for each method to the Branch Manager; and
	 Perform QA/QC audits, provide blind check samples and monitor results.

Position/Key Personnel

Responsibilities

Management Information Systems Department Damon Abbott

- Establish and maintain the computer systems, the network, and the LIMS (Laboratory Information Management System);
- Provide data deliverables in U.S. EPA CLP format with hard copy and disc deliverable formats;
- Develop software necessary to meet other client's deliverable formats;
- Interpret U.S. EPA CLP Statements of Work Organics and Inorganics for requirements and deliverables;
- Interface analytical instruments' data systems with our IBM Token Ring Network and develop software required at the interface;
- Train department personnel in various data entry software packages purchased or developed internally;
- Provide guidance and direction to other departments where required or requested or as determined by Branch Manager;
- Provide support to QA/QC Coordinator where necessary to automate, archive and generate QC trends from historical records; and
- Provide support and guidance for entire laboratory operation, from sample receiving through the generation of data deliverable packages, to streamline and increase productivity through computer automation.

Position/Key Personnel	<u>Responsibilities</u>
Lab Supervision C. Oliver (Organics) B. D. Hunt (GC/MS)	 Provide overall supervision of department/section operations;
Carl Causey (GC) Gary St Pere (Inorganics)	 Implement procedures consistent with the generation of legally defensible data;
	 Provide QC activities consistent with the branch QC procedures;
	 Provide additional QC activities, as needed, which are consistent with the Branch QA philosophies;
	 Provide final report review before releasing;
	 Provide work assignments to departmental personnel;
	 Provide analytical job training and cross-training within the department and between departments where applicable and warranted;
	- Provide corrective action for deficiencies;
	 Provide quarterly instrument detection limit (IDL) studies;
	- Maintain maintenance logs on all instruments;
	- Maintain all standards logs;
	 Provide leadership and management philosophies consistent with those of the Branch; and
	- Provide legally defensible data from each department.

LENL KEY PERSONNEL AND RESPONSIBILITIES

Position/Key Personnel

Responsibilities

Sample Receiving and Shipment Sharon Taber

- Provide sample control via entry of all parameters to be analyzed per sample in LIMS when sample arrives;
- Provide chain-of-custody receipt of samples externally, and internal from our walk-in cooler to analysts and prep;
- Provide sample analysis report daily for all samples in-house;
- Provide prepared sample bottles in refrigerated shippers;
- Provide for sample disposal/return to sender; and
- Keep track of supplies, order when needed, bill clients for those used.

LENL SAMPLE FLOW AND DOCUMENTATION CHART

Samı	ole Flow	Documentation					
Α.	Incoming Samples	Chain of CustodyRequest for AnalysisWork Authorization					
В.	Log-In and Assignment	Master LogbookTest Entry Form					
C.	Sample Handling and QC (1) Organics Prep Lab	 Extraction Record Percent Moisture Standards Prep Logs QC Sample Logs 					
	(2) GC Lab	 Chromatograms Bench Sheets Condition Reports Linearity Checks Surrogate Recovery Logs Blank, Standard and Spike Results Standards Logbook 					
	(3) GC/MS Lab	 Instrument Maintenance Logs Sample Logbook Quantitation Reports Chromatograms Initial Calibration Forms Continuing Calibration Checks Surrogate Recovery Logs Blank, Standard and Spike Results Computer Streamer Tapes Instrument Maintenance Logs 					
	(4) Metals Lab	 Digestion Log Bench Sheets Blank, Standard and Spike Results Percent Moisture Standards Prep Logs QC Sample Logs 					
D.	Reporting	Transmittal LettersTest Data ReportsGC/MS Data Reports					
E.	Sample Disposal	- Internal Sample Disposal Forms					

LENL-PENSACOLA 187202 LABWORKS SAMPLE/DATA FLOW **SAMPLE RECEIPT** INPUT/OUTPUT COMPUTER **ENTER LOCATION CODE IF** DATA ENTRY/ PRESENT OR CREATE IF NEW **PROCESSING** VERIFY CHAIN OF CUSTODY, WORK AUTH., CONTACT FORM, AND DATA **CONTACT FORM** SCREEN FOR AGREEMENT WORK AUTH. FORWARD TO CHAIN OF CUSTODY **OPERATIONS MANAGER** FOR RESOLUTION DOES NO INFORMATION AGREE? IS WORK AUTHORIZATION SIGNED? YES INPUT/OUTPUT SAMPLE FOLDER COMPUTER **GENERATE ROUTING SHEETS** ROUT. SHT. - GC/MS DATA ENTRY/ AND FORWARD TO DEPT. SUPV. ROUT, SHT, - GC **PROCESSING FOR SIGNOUT** ROUT, SHT. - METALS **DEPT. ANALYSES SAMPLES** AND FORWARDS TO DEPT. SUPV. **FOR REVIEW** SCHEDULE REPEAT **ANALYSIS** DOES DATA MEET QC SPECIFICATIONS YES INST. RAW DATA RECORD RESULTS ON ROUTING SHEETS AND FORWARD TO OFFICE FOR DATA ENTRY (WITH RAW DATA) INPUT/OUTPUT COMPUTER **RESULTS ARE ENTERED DAILY AS** DATA ENTRY/ RECEIVED. ROUTING SHEETS WITH **PROCESSING RAW DATA ARE FORWARDED TO** SAMP, CONTROL FOR FILING INPUT/OUTPUT QA SUPERVISOR OR DEPARTMENT COMPUTER SUPERVISOR CHECKS ALL DATA ON DATA ENTRY/ THE VALIDATION QUEUE **PROCESSING** TAKE APPR. ACTION **AS NEEDED TO** CORRECT PROBLEM IS DATA IN THE QUEUE NO ACCEPTABLE ? FINAL INVOICE INVOICE QUEUE YES REPORT QUEUE **FINAL TEST REP** SECRETARY RUNS ALL IN REPORT QUEUE AND INVOICE QUEUE AND MAILS TO CLIENT **END** HF NOTE: SPECIAL DELIVERABLES, IF REQUESTED ARE ALSO GENERATED AFTER VALIDATION STEP IS COMPLETED

ANALYTICAL REQUEST FORM

187203

LAW ENVIRONMENTAL, INC. 7215 PINE FOREST ROAD PENSACOLA FLORIDA 32536 904/944-9772

To:			Attn.:							
Fro	m:(Branch/C	Company Name)								
СО	C Number:		Project Number:							
Dat	e Shipped:		Date Results Red	quested:						
_	Sample ID	Analysis Requested	Detection Limits Req.	Sample Type	Method					
L	_									
\vdash										
H			-							
r										
L										
-										
L			 							
_										
_										
-										
Ļ										
\vdash										
F										

Comments:

HF



INORGANIC SAMPLE CHECKOUT FORM

Page ____ of ____

Contract:			Case:		SAS :	: SDG :							
ocation:				Calib. Ref:		Form Initiated:/							
_		СН	ECKED OUT			CHECKED IN							
Sample Types	Date	Time	Removal Purposes	Removed By	Custodian Approval	Date	Time	Returned By	Custodian Approval	Complete Y/N			
S/D	_		_	1	-				_				
			_						ļ				
_				_									
				 									
				1									
_				<u> </u>									
				-	<u> </u>			-					
			_						_				
	-			1		-	_						
Sample Ty		ole D = Digest											
						000 * * * *				* * * * *			
A - S	<u>Prepar</u> iolide	ation		F - Flame	Analysis AAS			<u>Disposition</u> L - Storage					
A - Solids B - AA/ICP Digestion				G - ICP				M - Submission					
	FAAS Digestio	n		H - GFAAS	FAAS N - Disposal								
D-H E-C	lg Digestion			I - Hg									
E-0	л пе г			J - CN K - Other									
				* * SAMPLES	CONSTITUTING	THIS BATCH *							
Matrix:			L	evel:			# of Phy	rsical Samples:					
_	Sample ID				Sample ID			Sample ID					
1 –				8			15	15					
2 -				9 ——		_	16	16					
3 -	·			10	_		17	17					
4 _				11			18						
5 _				12			19	19					
6 -				13 —			_ 20	20 ———					
7 -				14 —			_ 21	21					

ORGANIC SAMPLE CHECKOUT FORM

Contract:			Case:		SAS:			SDG:					
						Form initiated:/							
	CHECKED OUT CHECKED IN												
Sample Types S/X	Date	Time	Removal Purpose	Removed By	Custodian Approval	Date	Time	Returned By	Custodian Approval	Complete Y/N			
				 		_							
		<u> </u>	-	 	 	_							
-		<u> </u> 			-								
		<u> </u>						 	<u> </u>				
			1	 						 -			
								_	_				
				<u> </u>	<u> </u>			<u> </u>					
										 			
	<u> </u>		<u> </u>	 			<u> </u>						
Secondo 7	/pes S≃Samı			<u> </u>			<u> </u>	<u> </u>	<u></u>	<u>[</u>			
Semple 1)	/pes s=sami	94 A = EXITAC			D								
	Prepar		1		Removal Purpo Analysis	80			Disposition				
A-0	H / % Moisture	40541		G - Screen			}	•	2145AA WAT				
i	extraction of SV	,	1	H - VOA G) ,	M - Storage					
C-E	Extraction of PE	ST/PCB		1 - SV GC/I	MS			N - Submission					
f	Clean-up of SV		ĺ		CB GC/ECD			O - Disposal					
ì	lean-up of PES	ST/PCB	ļ		Confirmation								
F-C	Other		}	L - Other			ļ						
	• • • • • •		* * * * * *			• • • • •				* * * * *			
			• • • • • •	* * SAMPLES	CONSTITUTING	THIS BATCH *		• • • • •					
Matrix:			L(evel:			# of Ph	ysical Samples:					
	Samp	ole ID	į		Sample ID			Sample ID					
1 -				8		_	15	15					
2 -				9 —	_	-	16	16					
з -				10				_					
4 _	-			11									
5 –				12			19						
6 -				13			20						
			1										

QA/QC CORRECTIVE ACTION REQUEST FORM

REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:
DATA/DATES IMPACTED: DESCRIPTION: REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION: ACTION:	DATE:	DUE DATE:
DATA/DATES IMPACTED: DESCRIPTION: REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION: ACTION:	DATE:	DUE DATE:
REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:
REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:
REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:
REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:
REQUIRED ACTION: DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:DUE DATE:
DETERMINATION ASSIGNED TO: IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:
ASSIGNED TO:	DATE:	DUE DATE:
IMPLEMENTATION ASSIGNED TO: ACTION:	DATE:	DUE DATE:
ASSIGNED TO:		
ACTION:		
		· · · · · · · · · · · · · · · · · · ·
EFFECTIVENESS EVALUATION (S):		-
ASSIGNED TO:	DATE:	DUE DATE:
ASSIGNED TO:		
Addid to.	DATE	DOE DATE
CC: QA OFFICER		_
OPERATIONS MANAGER:		
GROUP LEADER:		
GROUP LEADER:		
SECTION SUPERVISOR:		

								-				Ŧ
		Ä.										
!	 } 	DATE										
	 1	DUE										j
!	11 14 16	CORRECTIVE ACTION DESCRIP										
<u> </u>	1	į	1	T								
IDEN	 	DATE INT.										
2		OUE										ĺ
DEFICIENT	 	CORRECTIVE ACTION DESCRIPTION										
SUMMARY:	S	BRIEF DESCRIPTION										
	STA	ORIGINATOR INT. & DATE					:					

APPENDIX B-2

QUALITY ASSURANCE PROGRAMS

Transglobal Environmental Geochemistry, (TEG)
Austin, Texas



TRANSGLOBAL

ENVIRONMENTAL

GEOCHEMISTRY, TEXAS

December 16, 1993

Ms. Darice Kurtzer Project Manager LAW ENVIRONMENTAL & ENGINEERING

SUBJECT: QUALITY CONTROL LINITS FOR 8020/8015 & LEAD FOR CARSWELL AFB PROJECT

Dear Ms. Kurtzer:

The following pages are the information that you requested. The control limits based on the historical data obtained from our mobile labs. If you have any questions regarding this data, please feel free to contact me.

Sincerely

Dr. Dernsing Luu President TEG/Texas

DL/ml

1. BTEX:

a. Quality Control Check (QCC)

Concentration (mg/L)	UCL.*	LCL*
1.00	1.15	0.85

UCL: Upper Control Limits LCL: Lower Control Limits

b. Method Detection Limits (5020/8020)

0.01 mg/L (Benzene, Toluene, & Ethylbenzene)
0.03 mg/L (Xylenes)

c. Matrxi Spike and Matrix Spike Duplicate

Spiked Concentration (mg/L)

1.00

Normal Range of & Recovery

80 - 120

Acceptable & Relative Deviation

± 25

187211

2. TPH (Modiifed 8015):

Gasoline:

a. Quality Control Check (QCC)

 Concentration (mg/L)
 UCL*
 LCL*

 200
 230
 170

UCL: Upper Control Limits LCL: Lower Control Limits

b. Method Detection Limits

1.0 mg/L

c. Matrxi Spike and Matrix Spike Duplicate

Spiked Concentration (mg/L)

200

Normal Range of & Recovery

75 - 125

Acceptable % Relative Deviation

± 27

Diesel:

a. Quality Control Check (QCC)

Concentration (mg/L)	UCL*	<u>LCL*</u>
500	565	435

UCL: Upper Control Limits LCL: Lower Control Limits

b. Method Detection Limits

2 mg/L

c. Matrxi Spike and Matrix Spike Duplicate

Spiked Concentration (mg/L)

500

Normal Range of *Recovery

85 - 115

Acceptable & Relative Deviation

<u>+</u> 25

- 3. Total Lead (3051/7421):
 - a. Quality Control Check (QCC)

1.0 UCL* LCL*

UCL: Upper Control Limits LCL: Lower Control Limits

b. Method Detection Limits

0.1 mg/L

c. Matrxi Spike and Matrix Spike Duplicate

Spiked Concentration (mg/L)

1.0

Normal Range of %Recovery

65 - 135

Acceptable & Relative Deviation

± 28

187214



Transglobal Environmental Geochemistry, Texas

December 16, 1993

Ms. Darice Kurtzer Project Manager LAW ENVIRONMENTAL & ENGINEERING

SUBJECT: QUALITY CONTROL LIMITS FOR 8020/8015 & LEAD FOR CARSWELL AFB PROJECT

Dear Ms. Kurtzer:

The following pages are the information that you requested. The control limits based on the historical data obtained from our mobile labs. If you have any questions regarding this data, please feel free to contact me.

Sincerely

Dr. Derhsing Luu President TEG/Texas

DL/ml

187215

1. BTEX:

a. Quality Control Check (QCC)

Concentration (mg/L)	UCL*	<u>LCL*</u>
1.00	1.15	0.85

UCL: Upper Control Limits LCL: Lower Control Limits

b. Method Detection Limits (5020/8020)

0.01 mg/L (Benzene, Toluene, & Ethylbenzene)
0.03 mg/L (Xylenes)

c. Matrxi Spike and Matrix Spike Duplicate

Spiked Concentration (mq/L)

1.00

Normal Range of Recovery

80 - 120

Acceptable % Relative Deviation

± 25



TRANSGLOBAL ENVIRONMENTAL GEOCHEMISTRY, TEXAS

TRANSGLOBAL ENVIRONMENTAL **GEOSAMPLING**

STRATAPROBE_{TM}

METHODOLOGY & REFERENCES

TABLE OF CONTENTS

- I STRATAPROBE_{TM}
- II SOIL VAPOR SAMPLING PROCEDURES
 - A) SOIL VAPOR COLLECTION

Probe Construction
Probe Insertion
Fig. A
Gas Sampling
Flushing & Decontamination Procedures
Access to Difficult Sites
Abandonment Procedures

- B) MULTI-DEPTH (Nested) VAPOR WELLS
 Well Construction
 Fig. B
 Tube Insertion
- C) SOIL VAPOR ANALYSIS
 Aromatic Hydrocarbons
 Chlorinated & Halogenated Hydrocarbons
 Total Petroleum Hydrocarbons
 Fixed/Biogenic Gases
 Nominal Throughput
 QA/QC Procedures
- D) DATA HANDLING & PROCESSING
- E) VAPOR PERMEABILITY STUDIES
- III SOIL & GROUNDWATER SAMPLING PROCEDURES
 - A) DISCRETE SOIL SAMPLER
 Sampler Construction
 Probe Insertion & Sample Recovery
 Decontamination Procedures
 - B) WATER SAMPLER
 Water Sampler Construction
 Water Sample Recovery
- IV RESUMES OF LEAD PERSONNEL
- V GEOSAMPLING REFERENCES
- VI GEOSAMPLING PROJECT DESCRIPTIONS
- VII LIST OF TRANSGLOBAL OFFICES / AFFILIATES

I STRATAPROBETM

The STRATAPROBETM is the basic element of a sophisticated hudraulic drive-point system developed by TEGeoSampling. TEG's STRATAPROBETM is designed to perform discrete soil sampling, groundwater sampling, soil vapor sampling, and the installation of multi-depth vapor wells for the Environmental Industry. The STRATAPROBETM is one of the most versatile units of its kind, capable of driving an assortment of sampling tools to depths of 50 feet, or greater, in most soil formations.

The STRATAPROBETM carrier vehicle is a four-wheel-drive, one ton truck, equipped with a reliable diesel-powered, PTO hydraulic system. The powerful, rear mounted unit is fitted with dual rams and a hydraulic hammer. High frequency impact energy coupled with 5000 pounds of static reaction weight and a 15000 pound pull-back capacity, provide ample force to overcome the most demanding geologic conditions. The 5' working stroke of the machine enables efficient penetration and retrieval of the sampling tools. Additionally, the low, 12' mast allows for operations within buildings, and the fully articulated boom provides a full range of positioning and directional boring capabilities. The custom-designed drive rods are constructed of high strength 11/2" threaded steel tubing. The STRATAPROBETM is a low profile unit that is fully enclosed in the utility bed of the truck while in transit. TEGeoSampling also offers hand-held, portable equipment for sampling in locations where truck access is not possible.

STRATAPROBE_{TM} SYSTEM

STRATAPROBETM CARRIER VEHICLE

Ford F350 Utility Truck

SPECIFICATIONS

- * Four Wheel Drive
- * All Terrain
- * Dimensions: 7.5' x 21'
- * Low Profile Bed
- Diesel Powered PTO

ACCESSORY EQUIPMENT

- * Steam Cleaner
- Concrete Corer
- Decontamination Area
- * Sample Prep & Waste Staging Area

STRATAPROBE_{TM} DRIVE POINT UNIT

Rear-Mounted Hydraulic Unit

SPECIFICATIONS

- * Dual 5' Stroke Hydraulics
- * High Frequency 100psi Impact Hammer
- * 5000lb Static Drive Weight
- * 15000lb Pull-Back Capacity
- * Custom Drive Rods: 5' Threaded, High-Strength Steel Tubing
- * 25° Directional Boring with Articulated Boom

II SOIL VAPOR SAMPLING PROCEDURES

A) SOIL VAPOR COLLECTION

TEG's experienced field personnel, equipped with hydraulic & vibrational insertion and recovery equipment, concentrate solely on vapor collection. TEG's probe design allows insertion/recovery using either heavy duty, 4-wheel drive truck mounted $STRATAPROBE_{TM}$ equipment or portable (vehicle-independent) equipment. No change-over time is required between the two methods, meaning both methods can be deployed simultaneously, depending upon site access. Our soil vapor probe design and methodology have been fully approved and endorsed by the California EPA and the Los Angeles Regional Water Quality Control Board (LA-RWQCB). The TEG system has been used on state lead projects for the TWC (references follow).

TEG's probe design has only 5cc dead-volume within an inert tube, which affords the following advantages:

- * Minimal vapor need be withdrawn, thereby eliminating the need for vacuum pumps.
- * Full length inert tubing prevents atmospheric contamination from leakage into probe body. This is a crucial consideration in arid environments.
- * Soil vapor does not contact the metal probe body. Thus, decon procedures are fast and simple.

Probe Construction

TEG's soil vapor probes are constructed of 7/8" or 1 1/2" OD ($STRATAPROBE_{TM}$) stainless steel, equipped with a hardened, reverse-thread steel tip. Nominal lengths are 5', although additional lengths may be used. An inert 1/8 inch nylaflow tube runs down the center of the probe to the sampling ports beneath the tip (fig A).

Probe Insertion

The probe is driven into the ground by the force of 5,000 lb of static weight and a high frequency hydraulic hammer with the $STRATAPROBE_{TM}$ system, or by a portable electric rotary hammer. Once inserted to the desired depth, the probe is rotated 3 to 5 times in a clockwise direction, which opens the tip and exposes the vapor sampling ports. This design prevents clogging of the sampling ports and cross-contamination from soils during insertion.

Field Collection Log

The field technician completes a logsheet summarizing time of sampling event, depth of pene-tration or refusal, which probe is used on each sampling location, when tubing is replaced, any visual contamination on the probe, OVM readings as applicable, and any other unusual occurences at a particular sampling location.

Gas Sampling

Soil vapor is withdrawn from the nylaflow tubing using a syringe connected via an on-off valve. The first 40cc of gas is discarded to flush out the dead volume of the tubing and fill with insitu soil vapor. The next 20cc of gas are drawn in a syringe, plugged, and immediately transferred to the mobile lab for analysis within minutes of collection. Additional soil vapor volumes may be collected and stored in gas-tight containers as desired.

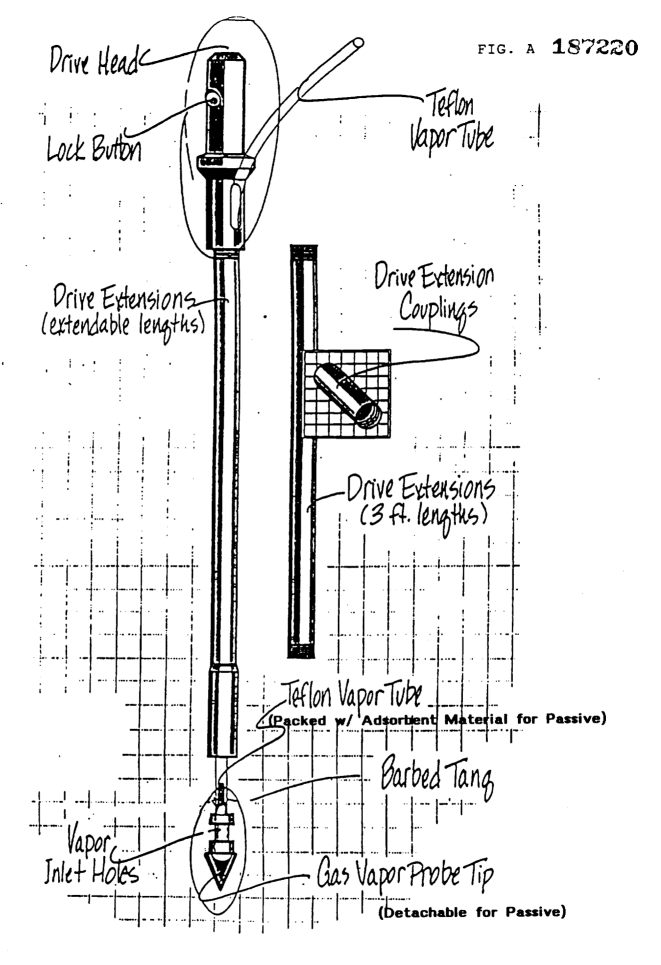


Diagram of TEG's Soil Gas Sampling Probe

Flushing & Decontamination Procedures

To minimize the potential for cross-contamination between sites, all probe parts are cleaned of excess dirt and moisture prior to insertion. The nylaflow tubing and sampling ports are flushed with hundreds of cc's of ambient air or inert gas between samples. If dirt, water or any material is observed in the tubing, it is discarded and replaced with new nylaflow. If concentrations greater than 100ppmv are detected for any compound (except methane), the tubing is replaced. Sampling syringes are opened and exposed to outside air on a clean surface to allow any volatiles to escape after each use. If concentrations greater than 100ppmv are detected for any compound (except methane) the syringe is discarded, unless glass syringes are used.

Access to Difficult Sites

TEG's probes may be deployed either with a truck-mounted system or by hand. The truck mounted $STRATAPROBE_{TM}$ system is deployed on a 4-wheel drive truck, thus enabling access to rough terrain sites. The portable unit may be used in and around buildings, inside tank farm fire walls, or in environmentally sensitive areas.

Abandonment Procedures

With only a 3/4" to 1 1/2" hole and no tailings remaining after recovering the probe, the hole is typically backfilled and grouted and the pavement or slab is patched accordingly.

B) MULTI-DEPTH (Nested) VAPOR WELLS

Well Construction

TEG's semi-permanent, multi-depth soil vapor wells are constructed with individual sample tubes set at multiple depths below the ground surface (Fig. B). An inert 1/8" polypropylene nylaflow tube extends to the desired depth and is connected to a machined, stainless steel tip containing multiple intake ports.

The 1/8" diameter nylaflow consists of one continuous length of tubing, which eliminates couplings or joints, and therefore, potential leakage. Using the 1/8" tubing also results in an extremely low dead volume of space within the tube and tip which must be purged prior to obtaining a sample. This low dead volume eliminates the need for purging large vloumes of air, which ensures sample integrity.

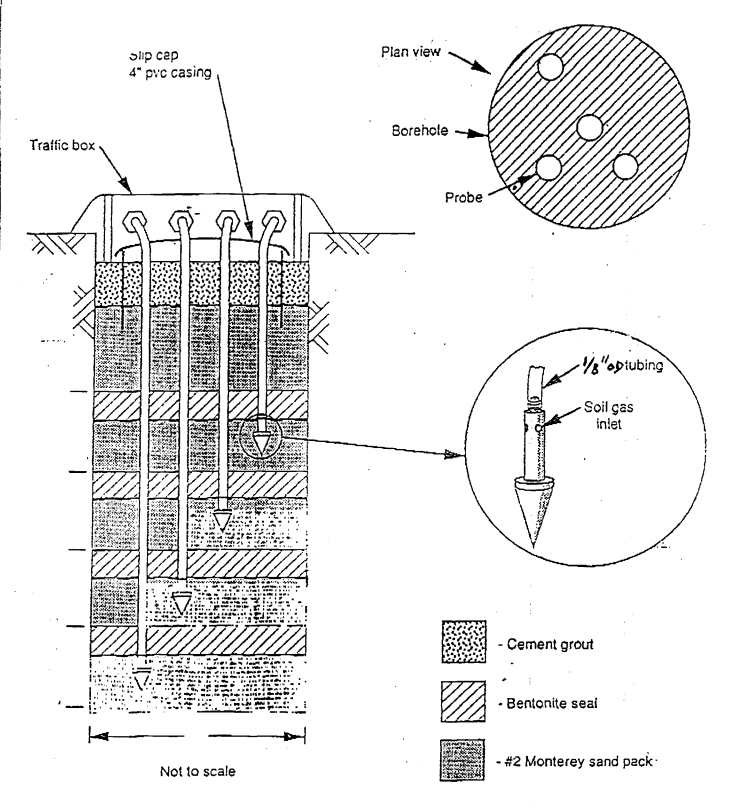
Tube Insertion

The 1/8' nylaflow tubing and tips can be inserted either using the $STRATAPROBE_{TM}$ to depths up to 50' or may be deployed at deeper depths in larger diameter borings. The $STRATAPROBE_{TM}$, equipped with a detachable, vapor slotted, drive point, is initially driven to the depth of the deepest vapor well. The probe is then withdrawn approximately 1', leaving the drive point with attached tubing in place. Clean sand is poured into the probe to provide a sand-pack around the vapor point, and bentonite grout is then pumped into the hole to insure a annular seal from the overlying strata. The probe is further removed and additional vapor wells are installed in the same manner, except the shallower wells are equipped with a screen instead of the vapor point.

Fig. B

NESTED VAPOR WELL

187222



C) SOIL VAPOR ANALYSIS

Soil Vapor is collected from each probe in a gas-tight syringe, brought to the TEG mobile laboratory, and analyzed immediately. Vapor will not be collected until the lab is ready for the sample, thus there is no lag time between gas collection and analysis (Very Important!).

Because there is some uncertainty about the types of contamination that might be present, it is *Imperative* that the analytical equipment be Sensitive enough and also Specific enough to differentiate between the different types of potential compounds. For example, FID and ECD detectors are sensitive enough, but used alone are not specific enough to differentiate between all of the aromatic and halogenated VOC's. To ensure proper identification and achieve adequate detection levels, a combination of 3 different types of detectors is often necessary: FID, Hall (ELCD), and PID.

Utilizing the Appropriate State of the Art Laboratory Grade analytical instrumentation With EPA Required Detectors, TEG's fully self-supported (power, water, etc.), CA DOHS Certified and TWC Approved Mobile Laboratories are equipped for the analyses of the following volatile organic compounds (VOC's) found in soil vapors:

Aromatic Hydrocarbons (BTEX, etc.), EPA modified 602, 8020

Gas Chromatograph with PID

Detection levels: 10-50 ppbv (0.1 ug/l) for each compound

Chlorinated & Halogenated Hydrocarbons (Solvents), EPA modified 601, 8010 Gas Chromatograph with ECD or Hall (ELCD) detector coupled with a 60m capillary column Detection levels: 10-50 ppbv (0.1 ug/l)

Total Petroleum Hydrocarbons (TPH), EPA modified 8015

Gas Chromatograph with FID

Detection level: 1ppmv

Fixed/Biogenic Gases (available at selected TEG offices)

Gas Chromatograph with combination TCD/FID

Detection levels:

fixed gases, 10ppm CO_2 , 10ppm CO, 10ppm methane, 1ppm

 0_2 , 0.5%

Nominal Throughput

Depending upon sampling depth and which analyses are required, 15 to 30 samples can be collected and analyzed in a 8 to 10 hour work day.

QA/QC Procedures

TRANSGLOBAL utilizes modified EPA protocols. Three point calibration curves will be run prior to the program and re-run when the daily continuing calibration falls outside of the limits (+/- 15%) as required by EPA protocols.

D) DATA HANDLING & PROCESSING

Data from each detector is processed by a chromatographic software system. The resulting values for each compound is entered into a spreadsheet file on a multi-tasking, 486/33 MHz PC. Print-outs of the data are available daily. An example print-out is enclosed.

The spreadsheet data is read into a contouring program (Surfer) and displayed graphically in both 2-D (contour) and 3-D (raised surface) projections. The lab is equipped with a color printer for onsite hardcopy output (examples follow). These projections are available on a real time basis.

Data is also be available to the client on computer disks in an ASCII format.

E) VAPOR PERMEABILITY STUDIES

Relative vapor permeabilities in the soil can easily be determined by attaching a small vacuum pump to the individual vapor probes and measuring the flow rate while maintaining a constant vacuum pressure of 20 to 40 inches of water. According to emperical results performed by our associates at sites in California, the radius of influence for this type of test is approximately 5'.

III SOIL & GROUNDWATER SAMPLING PROCEDURES

A) DISCRETE SOIL SAMPLER

The TEG $STRATAPROBE_{TM}$ obtains high quality discrete soil samples using a retractable piston, combined with a standard split-spoon sampler.

Sampler Construction

The $STRATAPROBE_{TM}$ split-spoon sampler is 12" long and has an outside diameter of 2.5". The internal piston assembly consists of a hardened steel tip, steel connecting rod and a teflon ring that prevents metal to metal contact between the piston and the sample liners. The split-spoon contains two or three 2" x 4" stainless steel, brass or aceatate retainer sleeves. During initial probe insertion, the piston is locked into place in the sampler body. The piston prevents the sampler from filling as it is advanced to depth.

Probe Insertion & Sample Recovery

The probe is driven into the ground using the heavy-duty rear-mounted hydraulic "direct push" rig. Once the target depth is reached, the piston is unlocked by a control rod and the sampler is driven 1' deeper, allowing the cored material to enter the sample tube as the piston retracts. The $STRATAPROBE_{TM}$ is then returned to the surface, where the sleeves are recovered from the split-spoon and sealed for delivery to TEG's "state approved" on-site mobile laboratory for immediate analysis.

Decontamination Procedures

EPA recommended protocol is the basis for all decontamination on TEG $STRATAPROBE_{TM}$ operations. The actual procedures are tailored for each particular project. The primary deciding factors used in determining the decontamination procedures are types of contaminants and the site conditions. The sampling tools are decontaminated and certified clean with the appropriate documentation provided.

B) WATER SAMPLER

Water Sampler Construction

The $STRATAPROBE_{TM}$ system water sampler consists of a detachable drive tip which is attached to a 24" retractable stainless steel well screen. The unit is encased in the lead probe tube. For water recovery the system can accomodate either a 20" bailer, capable of recovering 50ml of sample, or tubing that can be run into the probe after insertion and is connected to a peristalic pump at the surface.

Water Sampler Operation & Sample Recovery

After the water sampler is advanced into the water bearing zone the probe is withdrawn 2' to allow the retractable assembly to open to the formation. The water level is monitored by an electrical conductivity instrument and when the water has reached hydrostatic equlibrium within the well screen, the sample is ready to be taken. Depending upon the hydraulic conductivity of the formation and the constituents of concern, the sample will be taken with either the bailer or the post-run-tubing/peristalic pump system. At the surface the samples will be placed in VOA vials or other approved containers, chilled and perservatives added, as EPA protocol dictates.

IV RESUMES OF LEAD PERSONNEL

TEXAS & MEXICO

Mr. Rolf Woods, CPG, President TRANSGLOBAL SOUTH

Mr. Woods will run and/or supervise the day to day GeoSampling operation as well as market the services. In concert with these efforts, Mr. Woods will continue to promote mobile laboratory services for sister TEG operations on a commission basis.

Mr. Woods' professional career began as a well site geologist for UNOCAL where he worked in California, Thailand and Louisiana. During this time he was directly responsible for sampling and logging operations. Mr. Woods moved to Texas with Texas Oil & Gas Corporation in 1979 and became a consulting geologist in 1980. During the 1980's Mr. Woods generated drilling prospects and marketed these ideas to industry. In 1991 Mr. Woods performed soil vapor field operations on an exploration project for a consortium of independent oil and gas operators.

Since joining TEG in February, 1992 Mr. Woods has been instrumental in developing the mobile laboratory market in Texas. Due to his influence, TEG/TEXAS has also become a major player in the soil vapor surveying business. In the first 15 months of business Mr. Woods has developed a client list of over 600 companies in 27 states and Mexico.

Mr. Woods holds a BS degree from The University of Southern California in Geology and he is a Certified Petroleum Geologist.

SOUTHERN CALIFORNIA

Mr. Jeffrey S. Martineau, President TRANSGLOBAL ENVIRONMENTAL GEO-SAMPLING

Mr. Martineau is responsible for research and development of the geo-sampling equipment and technical support for field operations. He will also play a primary role in developing a marketing and pricing strategy for TRANSGLOBAL ENVIRONMENTAL GEO-SAMPLING.

Prior to joining the TEG family, Mr. Martineau was Western Operations Manager for TARGET ENVIRONMENTAL SERVICES. In this capacity, Mr. Martineau accumulated over 5 years of environmental site assessment experience and was the principal designer and builder of TARGET's sampling systems, which included deep soil vapor, soil and water sampling. He was instrumental in the construction of 2 innovative hydraulic drive point rigs. Mr. Martineau has completed more than 1000 geochemical surveys with special emphasis on difficult and varied sampling conditions throughout the United States.

Mr. Martineau holds a BS degree in Geography (Minor in Geology) from James Madison University.

SOUTHERN CALIFORNIA

Dr. Blayne Hartman, Ph.D., President TRANSGLOBAL ENVIRONMENTAL GEOCHEMISTRY

Dr. Hartman's daily responsibilities with respect to TEG include direction of laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review and method development, and direction of corporate development.

Dr. Hartman's academic research included a variety of chemical oceanography projects including: petroleum tracing using stable and radiogenic compounds as tracers; process controls across the air-water interface; and gas chromatography of dissolved gases in water and sediments.

Prior to founding Transglobal Exploration, Dr. Hartman was chief marine geochemist with InterOcean Systems where his responsibilities included direction of marine sediment and bottom-water geochemical exploration programs. As a research geochemist with Unocal Research, Dr. Hartman was responsible for development of geochemical prospecting techniques in onshore and offshore applications. These research operations included extensive method development and use of a variety of gas chromatography. Other research included use of stable metal isotopes as correlation parameters for organic materials. As founder and president of Transglobal Exploration & Geoscience in 1988, Dr. Hartman has been personally responsible for directing geochemical technical development, equipment design and maintenance, applications development, and implementation of exploration programs, analyses, and reporting.

TEXAS

Dr. Derhsing Luu, Ph.D., President TRANSGLOBAL ENVIRONMENTAL GEOCHEMISTRY / TEXAS

Dr. Luu's daily responsibilities with respect to TEG/TEXAS include direction of laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review and method development.

Dr. Luu's academic research included a variety of novel automated analytical instruments development including: methanol analyzer and hydrocarbon analyzer. Prior to founding TEG/Texas, Dr. Luu was R&D senior scientist with Global Geochemistry Corp. where his responsibility included development of methanol analyzer for CARB (California Air Resource Board) and hydrocarbon analyzer for Schlumberger. As a senior chemist for Air Toxic Lab with ENSR, Dr. Luu was responsible for daily laboratory operations and method development for air toxic laboratory. As an environmental laboratory manager and corporate QA officer with TRI (Texas Research Institute), Dr. Luu was responsible for supervising, performing GC/MS analysis, training, method developing for indoor air monitoring, and devising corporate QA/QC policy.

WASHINGTON

Michael Korosec, M.S., President TRANSGLOBAL ENVIRONMENTAL GEOCHEMISTRY / NW

Mr. Korosec's daily responsibilities with respect to TEG Northwest include direction of all laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review, marketing, and contract negotiations.

While with TEG, Korosec has developed unique preparation methods and analytical techniques for on-site analysis of PAH's, PCB's and chlorinated pesticides in soils, and PAH's and BTEX in tars.

Korosec's academic research included studies of the chemical and physical controls on the transport of dissolved nutrients across the sediment-water interface, using UV spectrophotometry and gas chromatography.

Prior to founding TEG/Northwest, Korosec was in charge of the state of Washington, Division of Geology and Earth Resources' Geothermal Exploration Program. The work included the development of a water analysis laboratory for the determination of dissolved cat ions, an ions and trace metals in thermal and mineral waters. Instrumentation included AA spectrophotometer, UV spectrophotometer, mercury analyzer and specific ion meters. Additional work included drilling programs for temperature gradient and heat-flow studies, whole rock geochemistry, age dating, and geologic mapping. As program manager, Korosec was responsible for all contracting, subcontracting and reporting to the U.S. Department of Energy.

NORTHERN CALIFORNIA

Mark Jerpbak, M.S., R.G., President TRANSGLOBAL ENVIRONMENTAL GEOCHEMISTRY / SACRAMENTO

Mr. Jerpbak's daily responsibilities with respect to TEG/Sacramento includes direction of all laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review, marketing, and contract negotiations.

Since 1981, Mr. Jerpbak has worked with several companies and agencies in geology and geochemistry in the United States and overseas, including: Ascension Island, Papau New Guinea and Japan. His work has included gas chromatography in the petroleum industry, gas and vapor analyses and wet chemical titration in the geothermal industry, injection tests in active geothermal areas, and gas chromatography by EPA methods in the environmental industry. He has worked extensively in the field and in the mobile lab environment.

Mr. Jerpbak holds a M.S. degree from the University of Iowa in geology and geophysics. He is a California State Registered Geologist and also consults for the Yolo County Department of Public Health, and Environmental Health Services in California.

V GEOSAMPLING REFERENCES

TRANSGLOBAL'S soil vapor sampling and analytical methodology have been fully approved by the LA County Regional Water Quality Control Board (RWQCB). TRANSGLOBAL has performed state lead soil vapor projects for the TWC. Selected references with contact numbers (we encourage you to contact them) for specific projects are:

CLIENT	CONTACT	PHONE #
AIRCRAFT STAMPING	Mr. Mike Nolan	818-443-2713
ALTON GEOSCIENCE	Mr. Ron Kofron	619-587-0682
CALIFORNIA ENV.	Mr. Charles Buckley	818-991-1542
CHEM WASTE MGMT	Ms. Sue Goss	714-826-0604
CERES ENVIRONMENTAL	Mr. Nick Patz	310-907-4088
COASTAL REMEDIATION	Mr. John Thomas	713-297-1708
DAMES & MOORE (Houston)	Mr. Jeff Borem	713-688-4541
DAMES & MOORE (LA)	Ms. Deborah Stott	213-683-1560
DAMES & MOORE (Tustin)	Mr. Essi Essamali	714-433-2000
DAMES & MOORE (San Ber.)	Mr. Brian Wynne	714-381-2004
DuPONT ENVIRON. REM. SERVICES	Mr. John Leetham	713-586-5600
ENSAFE	Mr. Ted Blahnik	214-642-1665
HARDING LAWSON ASSOC.	Mr. Dale Tischmak	303-292- <i>5</i> 365
W.W. IRWIN	Mr. David A. Williams	714-699-6120
IT CORPORATION (Irvine)	Ms. Mary Parker	714-660- <i>5</i> 466
IT CORPORATION (SD)	Mr. Peter Merz	619-554-0510
JACOBS ENGINEERING	Ms. Riz Sarmiento	818- <i>5</i> 68-7001
KLEINFELDER	Mr. Ed Trosper	310-860-5559
LOCKMAN & ASSOC	Mr. Razmik Gozalian	213-724-0250
MICHAEL BRANDMAN ASSOC	Ms. Susan Kline	619-260-1800
NINYO & MOORE	Mr. Steve Geyer	619-457-0400
OGDEN ENVIRONMENTAL	Mr. Eric Wetzstein	619-458-9044
MALCOLM PIRNIE	Mr. Dave Morley	713-840-1511
REIDEL ENVIRONMENTAL	Mr. Joe Koutski	310-327-4428
ROBERT PRATER ASSOC	Mr. Larry Jansen	619-453-5605
SEACOR	Mr. John Wainwright	714-335-6116
SIMON HYDROSEARCH	Mr. Roy Marroquin	714-891-7446
TETRA TECH	Mr. J.Ř. Hollingsworth	619-450-0365
TRAIL CHEMICAL	Mr. Bill Peters	818-442-4140
WEST COAST ENVIRONMENTAL	Ms. Carol Means	805-644-7976
WOODWARD-CLYDE	Mr. Tom Zep	714-835-6886
CA-RWQCB (LA)	Mr. Phil Chandler	213-266-7500
0.1 1 (0.2 (1.1)	Mr. Craig Christmann	213-266-7531
		

VI GEOSAMPLING PROJECT DESCRIPTIONS

Facility: Vought Aircraft, NWIRP, Dallas

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 600

Required Time to Complete Job: 7 weeks Consultant: EnSafe (Ted Blahnik, 214-642-1665)

Facility: Beale AFB, Sacramento

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 300

Required Time to Complete Job: 3 weeks

Consultant: Law Environmental (James Kozakowski, 916-649-2424)

Facility: Electronics Plant, Colorado Springs

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 75

Required Time to Complete Job: 4 days

Consultant: Harding Lawson (Dale Tischmak, 303-292-5365)

Facility: Schofield Army Barracks, Honolulu

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 400

Required Time to Complete Job: 25 days

Consultant: Harding Lawson (Kathrin Cain, 303-292-5365)

Facility: March AFB, Riverside

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 500

Required Time to Complete Job: 25 days

Consultant: Tetra Tech (Steve Williams, 714-660-5407)

Facility: 15 Service Station Sites in Rio Grande Valley Goals: Delineate contamination of aromatic hydrocarbons

Number of Sampling Points: 330

Required Time to Complete Job: 20 days

Consultant: Malcolm Pirnie (TWC state lead) (Dave Morley, 713-840-1511)

Facility: Tank Farm

Goals: Delineate contamination of aromatic hydrocarbons

Number of Sampling Points: 250

Required Time to Complete Job: 11 days

Consultant: Coastal Remediation (John Thomas, 713-297-1708)

Facility: Chemical Plant

Goals: Delineate contamination of aromatic hydrocarbons

Number of Sampling Points: 60

Required Time to Complete Job: 3 days

Consultant: DuPont Environ. Rem. Services (John Leetham, 713-586-5600)

Facility: Camp Pendleton Marine Corps Base

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 1500 Required Time to Complete Job: 43 days

Consultant: IT Corp. (Mary Parker, 714-660-5407)

Facility: Camp Naval Base, Guam

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 60

Required Time to Complete Job: 4 days

Consultant: Ogden Env. (David Bjostad, 619-458-9044))

Facility: Chemical Plant

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 110

Required Time to Complete Job: 7 days Consultant: Monsanto Chemical (Larry Adams)

Facility: Commercial Properties

Goals: Delineate contamination of aromatic and halogenated hydrocarbons

Number of Sampling Points: 225

Required Time to Complete Job: 15 days

Consultant: Dames & Moore (Nancy Darigo, 213-683-1560)

VII LIST OF TRANSGLOBAL OFFICES / AFFILIATES

TRANSGLOBAL SOUTH

408 University Place

Corpus Christi, TX 78412

512-991-7975 512-991-1078 Fax

Directors:

Mr. Rolf Woods, CPG

Units:

1 STRATAPROBE

Personnel:

TEG Labs/GeoSampling

432 Cedros Avenue

Solana Beach, CA 92075

619-793-0401 619-793-0404 Fax

Director:

Dr. Blayne Hartman

Labs:

5 mobile

Units:

1 STRATAPROBE

Personnel: 15

TEG - TEXAS

10805 Metric Blvd.

Austin, Texas 78758

512-835-9299

512-835-4726 Fax

Directors:

Dr. Derhsing Luu

Labs:

1 Fixed Base, 2 Mobile

Personnel:

4

TEG - Pacific Northwest

7110 38th Drive, S.E.

Lacey, WA 98503

206-459-4670

206-459-3432 Fax

Director.

Mr. Michael Korosek.

Labs:

3 mobile

Personnel:

TEG - Hawaii

770 Mokapu Road

Kailua. HÍ 96734

808-254-0046

808-254-0243 Fax

Director:

Mr. Tim Fitzpatrick

Labs:

2 mobile

Personnel:

TEG - Northern California

P.O. Box 162580

Sacramento, CA 95816

916-736-3233 916-452-5806 Fax

Director

Mr. Mark Jerpak, M.S.

2 mobile Labs:

Personnel: 4

TEG - Georgia

5064 Laurel Bridge Drive

Smyrna, GA 30082

404-433-8195

Director.

Mr.Mark Hankinson

Labs:

1 mobile

Personnel 3

COMPREHENSIVE QUALITY ASSURANCE PLAN (COMPQAP)

Prepared by

TRANSGLOBAL ENVIRONMENTAL GEOCHEMISTRY, TEXAS

10805 METRIC BLVD.

AUSTIN, TEXAS 78758

(512) 835-9299

Derhsing Lyu, Ph.D.
President

Sallie Cordell
Vice-President

01/01/93

01/01/92

Date

Section 2.0
Revision 1
Date 1/1/93
Page 1 of 5
187234

TABLE OF CONTENTS

Secti	<u>on</u>	# of Pages	Revision Date
1.0	TITLE PAGE	1	1-1-93
2.0	TABLE OF CONTENTS	5	1-1-93
3.0	STATEMENT OF POLICY	1	1-1-93
4.0	ORGANIZATION AND RESPONSIBILITY 4.1 Purpose 4.2 Corporate Profile 4.3 Key Personnel 4.4 Responsibilities	6	1-1-93
5.0	QUALITY ASSURANCE OBJECTIVES	5	1-1-93
6.0	SAMPLING PROCEDURES 6.1 Sample Containers, Preservation, Holding 6.2 Cleaning 6.3 Waste Disposal	2	1-1-93
7.0	 SAMPLE CUSTODY PROCEDURES 7.1 Sample Custody Objectives 7.2 Field Custody 7.3 Laboratory Custody 7.4 Electronic Data Records 	11	1-1-93
8.0	ANALYTICAL PROCEDURES 8.1 Method References 8.2 EPA Methods 601 and 8010 8.3 EPA Methods 602 and 8020 8.4 EPA Method 8015 TPH 8.5 EPA Method 418.1 8.6 EPA Method 601/8100 8.7 EPA Method 608/8080 8.8 EPA Method 604/8040 8.9 Reagent Storage	18	1-1-93

Section		# of Pages	Revision Date	
9.0	CALIBRATION 9.1 Instrumentation List 9.2 Standard Receipt and Traceability 9.3 Standard Sources and Preparation 9.4 Instrumentation Calibration 9.4.1 Instrument Group Calibration 9.4.2 Initial Calibration and Linearity 9.4.3 QC Check Standards 9.4.4 Continuing Calibration	10	187235 1-1-93	
10.0	9.4.5 Equipment Monitoring PREVENTIVE MAINTENANCE PROCEDURES AND SCHEDULES 10.1 Routine Maintenance 10.2 Documentation 10.3 Contingency Plan	3	1-1-93	
11.0	QUALITY CONTROL CHECKS, PRECISION AND ACCURACY AND DETECTION LIMITS 11.1 Laboratory QC Checks 11.1.1 Equipment QC Checks 11.1.2 Chemistry QC Checks 11.1.3 Additional QC Checks 11.1.3 Additional QC Checks 11.2 Precision and Accuracy 11.3 Method Detection Limits and Practical Detection Limits	3	1-1-93	
12.0	DATA REDUCTION, VALIDATION AND REPORTING 12.1 Data Reduction 12.2 Data Validation 12.3 Data Reporting 12.4 Data Storage	1	1-1-93	
13.0	CORRECTIVE ACTION 13.1 Criteria 13.2 Responsible Individuals 13.3 Corrective Actions 13.4 External Sources 13.5 Notification of Personnel	3	1-1-93	

Section 2.0 Revision 1 Date 1/1/93 Page 3 of 5

Section			# of Pages	Revision Date	
					187236
14.0	PER	FORMANCI	E AND SYSTEM AUDITS	3	1-1-93
	14.1	General Re	quirements		
	14.2	System Au	dits	•	
		14.2.1	System Audit Scope		
		14.2.2	Internal Audit Scope		
		14.2.3	External System Audit		
	14.3	Performance	æ Audit		
		14.3.1	Performance Audit Scope		
		14.3.2	Internal Performance Audits	5	
		14.3.3	External Performance Audit	is .	
15.0	QUA	LITY ASSU	FRANCE REPORTS	1	1-1-93
16.0	Ms. S Dr. B Dr. D Mr. N	JMES OF Sheri W. Hard layne Hartma Jerhsing Luu, Michael Koros Mark Jerpbak	nn, Ph.D. Ph.D. sec, M.S.	11	1-1-93

Section 2.0 Revision 1 Date 1/1/93 Page 4 of 5

187237

FIGURES

<u>Figur</u>	<u>re</u>	<u>Sec</u>	ction / Page	Revision Date
4.1	Corporate Organizational Chart		4 / 1	1-1-93
4.2	Sample of Data Flow Chart	4 / 4	1-1-93	
7-1	Chain-of-Custody Record		7 / 4	1-1-93
7-2	Extraction Log		7 / 5	1-1-93
7-3	Analysis Log		7 / 6	1-1-93
7-4	Project / Report Checklist Form		7 / 7	1-1-93
9-1	Solvent and Standard Log-In Form	9 / 2	1-1-93	
9-2	Standard Preparation Log		9 / 3	1-1-93
9-3	Instrument Initial Calibration Log	9/6	1-1-93	
9-4	Continuing Calibration Form		9 / 7	1-1-93
9-5	Balance Calibration Log		9 / 9	1-1-93
9-6	Refrigerator Temperature Log		9 / 10	1-1-93
10-1	Repair Maintenance Log		10 / 3	1-1-93
14-1	Internal System Audit Form	14 / 3	1-1-93	

Section 2.0
Revision 1
Date 1/1/93
Page 5 of 5
187233

TABLES

Table	<u>es</u>		Section / Page	Revision Date
5-1	QA Objectives		5 / 2	1-1-93
6-1	Containers, Preservation and Holding Times	6 / 1	1-1-93	
8-1	Reagent and Standard Storage		8 / 18	1-1-93
9-1	Instrumentation List	9 / 1	1-1-93	
9-2	Instrument Calibration		9 / 5	1-1-93
10-1	Maintenance Activities		10 / 1	1-1-93
11-1	Equipment Monitoring Activities		11 / 1	1-1-93
11-2	Chemistry QC Checks		11 / 2	1-1-93
13-1	Summary of Corrective Action Procedures		13 / 2	1-1-93

187239

3.0 STATEMENT OF POLICY

This document has been prepared for the purpose of providing detailed information to regulatory agencies and clients regarding the quality assurance (QA) and quality control (QC) program which is utilized at Transglobal Environmental Geochemistry, Inc. (TEG). The format of this Comprehensive Quality Assurance Plan is generally based on United States Environmental Protection Agency (EPA) guidelines for preparation of Quality Assurance Program and Project Plans (QAMS-004/80 and QAMS-005/80) and the Quality Control portions of EPA Document SW 846, "Test Methods for Evaluating Solid Waste." Changes to those formats have been incorporated to expand this document into a non-project specific outline of TEG operating procedures.

The purpose of TEG's quality assurance program is to establish methods by which problems in the data acquisition process may be detected and corrected within a framework of statistical control. This approach is designed to result in reduction of measurement errors to within accepted standards with results being of acceptable and verifiable quality. The TEG quality assurance program is concerned with, but is not limited to, the following subjects:

- development and use of techniques commonly accepted as good laboratory practice;
- standard operating procedules used with consistency;
- adherence to protocol for specific analyses;
- continuity and consistency of personnel involved with all phases of the analytical program;
- consistent use and maintenance of equipment;
- application of proper calibration protocol and use of standards; and
- integration of senior technical and management personnel in supervision, evaluation, and implementation of the QA program.

By using the techniques outlined in this plan, the quality of the analytical process and resulting data is continually evaluated. In maintaining sound QA management practices, TEG is able to maintain its commitment to generating data of verifiable quality.

4.0 ORGANIZATION AND RESPONSIBILITY

187240

4.1 PURPOSE

Transglobal Environmental Geochemistry, Inc. (TEG) is seeking full and unconditional DER approval of laboratory services for on-site testing of soils, water, sediments, wastes, air, soil vapor, and air emissions for the EPA and DER methods described in Section 5.0 of this document.

4.2 CORPORATE PROFILE

Transglobal Environmental Geochemistry is a fully-owned subsidiary of Transglobal Exploration & Geoscience, Inc., a California Corporation based in Solana Beach, California. TEG has formed joint partnerships in five (5) additional locations outside of southern California as shown in Figure 4-1.

Figure 4-1. CORPORATE ORGANIZATION CHART

TRANSGLOBAL EXPLORATION & GEOSCIENCE, INC.

432 N. Cedros Avenue Solana Beach, CA 92075

TEG - SOLANA BEACH

432 N. Cedros Avenue Solana Beach, CA 92075 Blayne Hartman, Ph.D.

TEG - TEXAS

7013 Dallas Dr. Austin, TX 78729 Derhsing Luu, Ph.D.

TEG - NORTHWEST

7110 38th Dr. S.E. Lacey, WA 98503 Michael Korosec, M.S.

TEG - SACRAMENTO

3227 Second Ave. P.O. Box 162580 Sacramento, CA 95816 Mark Jerpbak, M.S.

Section 4 Revision 1 Date 1/1/93 Page 2 of 6

4.3 KEY PERSONNEL

At each location, all aspects are under the supervision of a CADOHS-certified director. This director is responsible for all aspects of laboratory operations, analytical methods, technical development, and supervision of analysis. The Director for each location is provided (Figure 4.1) and a brief description of these key personnel is provided in the following paragraphs.

SOUTHERN CALIFORNIA

Dr. Blayne Hartman, Ph.D.

Dr. Hartman's daily responsibilities with respect to TEG include direction of laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review and method development, and direction of corporate development. Additional discussion of responsibility is presented in Section 4.4.

Dr. Hartman's academic research included a variety of chemical oceanography projects including: petroleum tracing using stable and radiogenic compounds as tracers; process controls across the air-water interface; and gas chromatography of dissolved gases in water and sediments.

Prior to founding Transglobal Exploration, Dr. Hartman was chief marine geochemist with InterOcean Systems where his responsibilities included direction of marine sediment and bottom-water geochemical exploration programs. As a research geochemist with Unocal Research, Dr. Hartman was responsible for development of geochemical prospecting techniques in onshore and offshore applications. These research operations included extensive method development and use of a variety of gas chromatography. Other research included use of stable metal isotopes as correlation parameters for organic materials. As founder and president of Transglobal Exploration & Geoscience in 1988, Dr. Hartman has been personally responsible for directing geochemical technical development, equipment design and maintenance, applications development, and implementation of exploration programs, analyses, and reporting.

TEG - TEXAS

Dr. Derhsing Luu, Ph.D.

Dr. Luu's daily responsibilities with respect to TEG/Texas include direction of laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review and method development.

Dr. Luu's academic research included a variety of novel automated analytical instruments development including: methanol analyzer and hydrocarbon analyzer.

Section 4
Revision 1
Date 1/1/93
Page 3 of 6

Prior to founding TEG/Texas, Dr. Luu was R&D senior scientist with Global Geochemistry Corp. where his responsibility included development of methanol analyzer for CARB (California Air Resource Board) and hydrocarbon analyzer for Schlumberger. As a senior chemist for Air Toxic Lab with ENSR, Dr. Luu was responsible for daily laboratory operations and method development for air toxic laboratory. As an environmental laboratory manager and corporate QA officer with TRI (Texas Research Institute), Dr. Luu was responsible for supervising, performing GC/MS analysis, training, method developing for indoor air monitoring, and devising corporate QA/QC policy.

TEG - NORTHWEST

Michael Korosec, M.S.

Mr. Korosec's daily responsibilities with respect to TEG Northwest include direction of all laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review, marketing, and contract negotiations.

While with TEG, Korosec has developed unique preparation methods and analytical techniques for on-site analysis of PAH's, PCB's and chlorinated pesticides in soils, and PAH's and BTEX in tars.

Korosec's academic research included studies of the chemical and physical controls on the transport of dissolved nutrients across the sediment-water interface, using UV spectrophotometry and gas chromatography.

Prior to founding TEG/Northwest, Korosec was in charge of the state of Washington, Division of Geology and Earth Resources' Geothermal Exploration Program. The work included the development of a water analysis laboratory for the determination of dissolved cat ions, an ions and trace metals in thermal and mineral waters. Instrumentation included AA spectrophotometer, UV spectrophotometer, mercury analyzer and specific ion meters. Additional work included drilling programs for temperature gradient and heat-flow studies, whole rock geochemistry, age dating, and geologic mapping. As program manager, Korosec was responsible for all contracting, subcontracting and reporting to the U.S. Department of Energy.

TEG - SACRAMENTO

Mark Jerpbak, M.S., R.G.

Mr. Jerpbak's daily responsibilities with respect to TEG/Sacramento includes direction of all laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review, marketing, and contract negotiations.

Since 1981, Mr. Jerpbak has worked with several companies and agencies in geology and geochemistry in the United States and overseas, including: Ascension Island, Papau New

Section 4 Revision 1 Date 1/1/93 Page 4 of 6

Guinea and Japan. His work has included gas chromatography in the petroleum industry, gas and vapor analyses and wet chemical titration in the geothermal industry, injection tests in active geothermal areas, and gas chromatography by EPA methods in the environmental industry. He has worked extensively in the field and in the mobile lab environment.

Mr. Jerpbak holds a M.S. degree from the University of Iowa in geology and geophysics. He is a California State Registered Geologist and also consults for the Yolo County Department of Public Health, and Environmental Health Services in California.

4.4 RESPONSIBILITIES

The mobile laboratories based in each location are personally supervised by one of the laboratory directors with the assistance of trained and experienced technicians. It is the responsibility of each director to maintain the QA/QC protocol or SOP set forth in this document. In order to review adherence to these procedures, periodic meetings are established to review project data and related QA concerns. Additional discussion of responsibilities regarding each position is presented in following paragraphs. A sample and data flow chart is provided below.

Figure 4-2. SAMPLE OF DATA FLOW CHART

Sample Collection (Client)

Sample Tracking Sample Log-in (Lab Chemist) Sample Tracking and Storage (Lab Chemist)	Analysis (Lab Chemist)	Preliminary Report (Lab Chemist)
--	---------------------------	--

QA Review (QA Manager)

Final Report (Laboratory Director)

The responsibility for maintaining the QA/QC program is divided into separate designated tasks, each with differing requirements as follows:

Section 4
Revision 1
Date 1/1/93
Page 5 of 6

<u>Lab Chemist</u>. The lab chemist, or chemical analyst/technician, is responsible for implementing QA/QC procedures and record keeping in the preparation and analysis of samples, and for providing the initial review and interpretation of data within the framework of the QA/QC plan. In field situations, the lab chemist is also responsible for monitoring sampling activities, sample receipt, and sample control. Sample control duties include:

- sample receipt, unpacking, inspection, and verification;
- Chain-of-Custody Report completion;
- log-in of samples on proper tracking forms;
- placing samples in proper storage;
- notification of analysts and managers; and
- disposal or return of samples to client.

Analytical duties of the Lab Chemist include:

- daily maintenance, start-up and calibration of analytical equipment;
- daily monitoring of quality control protocol, such as refrigerator temperature calibration:
- preparation of standards for linearity checks;
- documentation of analyses, problems, QA, and maintenance of project files; and
- preparation of preliminary analytical report.

<u>QA Manager</u>. The QA Manager, is responsible for: establishing QA/QC procedures and specifications for each project; reviewing data and protocol; initiating and supervising audits; recommending appropriate corrective actions, and reporting to the directors. Specifically, the QA Manager is responsible for:

- data assembly, general review of project;
- verification of data completeness;
- verification of QA/QC compliance;
- verification of client requirements;
- review of preliminary report;
- preparation of QA report to include: technical difficulties, QA/QC results and conclusions; and
- implementation of the TEG QA program and technical training of personnel.

<u>Health and Safety Officer</u>. The Health and Safety (H&S) Officer at TEG is responsible for administrating the following duties:

- coordination of training programs in first aid, CPR, hazardous materials handling, emergency contingency planning, and right-to-know compliance:
- review and implementation of in-house and site-specific H&S Plans;

Section 4 Revision 1 Date 1/1/93 Page 6 of 6

- labeling and storage of hazardous materials;
- maintenance of Material Safety Data Sheets (MSDS) files;
- Resource Conservation Recovery Act small quantity generator record keeping;
 and
- H&S equipment ordering, maintenance, and record keeping.

Laboratory Director. It is the responsibility of the director to implement and maintain the overall quality assurance program at TEG. The director must meet regularly with the QA Manager to discuss general adherence to the program as well as specific QA problems or projects which require additional review and corrective action. Additional responsibilities of the Director include:

- preparation or review of SOPs and QA/QC protocol;
- implementation, updating and distribution of SOPs;
- training of key personnel;
- document control, security and confidentiality;
- technical application and development;
- resolution of QA and technical problems; and
- overall business management.

Section 5
Revision 1
Date 1/1/93
Page 1 of 7

5.0 QUALITY ASSURANCE OBJECTIVES

This section summarizes the quality assurance objectives for precision, accuracy, and method detection limit. These criteria are listed in Table 5-1 for each analytical parameter. The listed criteria are objectives for analysis of typical samples under nominal conditions and may not apply to samples in complex interfering matrices or to measurement of concentrations less than the listed detection limit for the method.

Precision, the measurement of mutual agreement among individual measurements of the same property under similar conditions, is expressed in terms of the percent maximum allowable relative difference, as standard deviation, within a set of replicate results and is assessed from duplicate and replicate analyses of the same sample. Accuracy goals are expressed as a range of acceptable recovery from the true value as assessed from standard deviation of reference samples and percent recoveries from matrix spikes and spike duplicates. Additional information regarding calculations for QA objectives may be found in Section 11.

TABLE 5-1 QUALITY ASSURANCE OBJECTIVES

COMPONENT	<u>MATRIX</u>	METHOD	PRECISION SRPD (Max.)	² ACCURACY <u>%Recovery</u>	MDL
PURGEABLE AROMATICS	Groundwater Surface Water Effluents	EPA 5030/602			us/L
Benzene Toluene Chlorobenzene Ethylbenzene 1,3-Dichlorobenzen 1,2 and 1,4-Dichlor m&p-Xylene o-Xylene	ıe	. · ·	75-110 15 15 15 15 15 15 15	0.30 75-110 75-110 75-110 75-110 75-110 75-110	0.30 0.30 0.30 0.30 0.30 0.30
PURGEABLE AROMATICS	Soils Sediments Solid Waste	EPA 3550/5030/8020			mg/kg
Benzene Toluene Chlorobenzene Ethylbenzene 1,3-Dichlorobenzen 1,2 and 1,4-Dichlor m&p-Xylene o-Xylene	ie	15	75-110 15 15 15 15 15 15	0.005 75-110 75-110 75-110 75-110 75-110 75-110	0.005 0.005 0.005 0.005 0.005 0.005
TOTAL RECOVERABLE PETROLEUM HYDROCARBONS	Water S	EPA 3510/418.1			me/L 5.0
TOTAL RECOVERABLE PETROLEUM HYDROCARBONS	Soils S	EPA 3550/418.1			mg/kg 0.5
TOTAL PETROLEUM HYDROCARBON	Groundwater Surface Water ** Effluents	EPA 3510/5030/8015 modified	15	65-110	<u>me/L</u> 0.5
TOTAL PETROLEUM HYDROCARBON	Soils ** Sediments Solid Waste	EPA 3550/8015 modified	15	65-110	mg/kg 5.0

Modifications are discussed in Section 8.3.

Listed MDLs are from published EPA methods or based on historical TEG data.

^{••} CA-DOHS modified EPA 8015 for fuels

^{1.} Refer to Section 11.2 for definition of Precision.

^{2.} Refer to Section 11.2 for definition of Accuracy, listed values from in-house data. Values also apply to surrogates.

Section 5
Revision 1
Date 1/1/93
Page 3 of 7

TABLE 5-1 (continued) QUALITY ASSURANCE OBJECTIVES

COMPONENT MATRIX	<u>METHOD</u>	'PRECISION %RPD (Max)	ACCURACY %Recovery*	MDL
PURGEABLE Groundwater HALOCARBONS Surface water Effluents				ue/L
Bromoform		15	13-159	0.3
Bromomethane		15	D-144	0.3
Carbon Tetrachloride		15	43-143	0.3
Chlorobenzene		15	38-150	0.3
Chloroethane		15	46-137	0.3
Chloroform		15	49-133	0.3
Chloromethane		15	D-193	0.3
Dibromochloromethane		15	24-191	0.3
1,3-Dichlorobenzene		15	7-187	0.3
2-and 1,4-Dichlorobenzene			15	42-1430.3
1,1-Dichloroethane		15	47-132	0.3
1,2-Dichloroethane		15	51-147	0.3
1,1 Dichloroethene		15	28-167	0.3
trau		15	38-155	0.3
1,2-Dichloropropane		15	44-156	0.3
cis-1,3-Dichloropropene		15	22-178	0.3
trans-1,3-Dichloropropene	15	22-178		0.3
Ethylene Dibromide		15	50-150	0.3
Methylene Chloride		15	25-162	0.3
1,1,2,2-Tetrachloroethane		15	8-184	0.3
Tetrachloroethane		15	26-162	0.3
1,1,1-Trichloroethane		15	41-138	0.3
1,1,2-Trichloroethane		15	39-136	0.3
Trichloroethane		15	35-146	0.3
Trichlorofluoromethane		15	21-156	0.3
Vinyl Chloride		15	28-163	0.3

Recovery limits taken from SW 846.

Listed MDLs are from published EPA methods or based on historical TEG data.

^{1.} Refer to Section 11.2 for definition of Precision.

^{2.} Refer to Section 11.2 for definition of Accuracy, listed values from in-house data. Values also apply to surrogates.

Section 5 Revision 1 Date 1/1/93 Page 4 of 7

TABLE 5-1 (continued) QUALITY ASSURANCE OBJECTIVES

PRECISION ACCURACY COMPONENT MATRIX	METHOD	%RPD (Max)	%Recovery	MDL
PURGEABLE Soils and HALOCARBONS Sediments	EPA 3550/5030/8010			mg/kg
Bromoform		15	60-125	0.005
Bromomethane		15	60-125	0.005
Carbon Tetrachloride		15	65-120	0.005
Chlorobenzene		15	65-120	0.005
Chloroethane		15	65-120	0.005
Chloroform		15	65-120	0.005
Chloromethane		15	65-120	0.005
Dibromochloromethane	,	15	65-120	0.005
1,3-Dichlorobenzene		15	65-120	0.005
1,2 and 1,4-Dichlorobenzene			15	65-1200.005
1,1-Dichloroethane		15	65-120	0.005
1,2-Dichloroethane		15	65-120	0.005
1,1-Dichloroethene		15	65-120	0.005
trans-1,2-Dichloroethene		· 15	60-125	0.005
1,2-Dichloropropane		15	60-125	0.005
cis-1,3-Dichloropropene		15	60-125	0.005
trans-1,3-Dichloropropene	15	60-125		0.005
Ethylene Dibromide				0.005
Methylene Chloride		15	60-125	0.005
1,1,2,2-Tetrachloroethane		15	60-125	0.005
Tetrachloroethane		15	60-125	0.005
1,1,1-Trichloroethane		15	60-125	0.005
1,1,2-Trichloroethane		15	60-125	0.005
Trichloroethane		15	60-125	0.005
Trichlorofluoromethane		15	60-125	0.005
Vinyl Chloride		15	50-125	0.005

Listed MDLs are from published EPA methods or based on historical TEO data.

1. Refer to Section 11.2 for definition of Precision.

Refer to Section 11.2 for definition of Accuracy, listed values from in-house data. Values also apply to surrogates.

Section 5 Revision 1 Date 1/1/93 Page 5 of 7

TABLE 5-1 (continued) QUALITY ASSURANCE OBJECTIVES

COMPONENT	<u>MATRIX</u>	METHOD	'PRECISION %RPD (Max)	² ACCURACY <u>%Recovery</u>	MDL
	Groundwater urface water Effluents	EPA 3510/610			ue/L
Acenapthene			20	65-125	10
Acenapthylene			20	65-125	10
Anthracene			20	65-125	10
Benzo(a)anthracene			20	65-125	20
Benzo(a)pyrene			20	65-125	20
Benzo(b)fluoranthene			20	65-125	20
Benzo(ghi)perylene			20	65-125	20
Benzo(k)fluoranthene			20	65-125	20
Chrysene			20	65-125	20
Dibenzo(a,h)anthracen	ne .		20	65-125	30
Fluoranthene			20	65-125	10
Fluorene			20	65-125	10
Indeno(1,2,3-cd)pyren	e		20	65-125	30
Napthalene Phenanthrene			20	65-125	10
Pyrene Pyrene			20	65-125	10
ryrene			20	65-125	10
	Soils and Sediments	EPA 3550/8100			mg/kg
Acenapthene			20	65-125	0.1
Acenapthylene			20	65-125	0.1
Anthracene			20	65-125	0.1
Benzo(a)anthracene			20	65-125	0.1
Benzo(a)pyrene			20	65-125	0.1
Benzo(b)fluoranthene			20	65-125	0.1
Benzo(ghi)perylene			20	65-125	0.1
Benzo(k)fluoranthene			20	65-125	0.1
Chrysene			20	65-125	0.1
Dibenzo(a,h)anthracene	e		20	65-125	0.5
Fluoranthene Fluorene			20	65-125	0.2
Indeno(1,2,3-cd)pyrene			20	65-125	0.2
Napthalene	•		20 20	65-125	0.5
Phenanthrene			20	65-125	0.2
Pyrene			20	65-125	0.2
1 71 cue			20	65-125	0.2

Listed MDLs are from published EPA methods or based on historical TEG data.

Refer to Section 11.2 for definition of Precision.

^{2.} Refer to Section 11.2 for definition of Accuracy, listed values from in-house data. Values also apply to surrogates.

Section 5 Revision 1 Date 1/1/93 Page 6 of 7

TABLE 5-1 (continued) QUALITY ASSURANCE OBJECTIVES

COMPONENT	MATRIX	METHOD	'PRECISION %RPD (Max)	² ACCURACY <u>%Recovery</u>	MDL
PHENOLS	Groundwater Surface water Effluents	EPA 3510/604			ue/L
2,3,4-Trichlorophe	enol		20	65-125	.01
2,4,6-Trichlorophe			20	65-125	.01
2,4-Dichloropheno	l		20	65-125	.01
2,4-Dimethylpheno	ol		20	65-125	.01
2,4-Dinitrophenol			20	65-125	.03
2,6-Dichloropheno	1		20	65-125	.03
2 - Methylphenol		20	65-125		.01
2 - Chlorophenol			20	65-125	.01
2 - Methyl 4,6 - D	initrophenol		20	65-125	.03
2 - Nitrophenol			20	65-125	.01
4 - Methylphenol		20	65-125		.03
4 - Chloro 3 - Met	hylphenol		20	65-125	.01
4 - Nitrophenol			20	65-125	.03
Pentachlorophenol			20	65-125	.01
Phenol			20	65-125	.01
PHENOLS	Soils and	EPA 3550/8040			
	Sediments				me/ke
2,3,4-Trichlorophe	nol		20	65-125	.2
2,4,6-Trichlorophe	nol		20	65-125	.2
2,4-Dichlorophenol			20	65-125	.2
2,4-Dimethylpheno	1		20	65-125	.2
2,4-Dinitrophenol			20	65-125	.5
2,6-Dichlorophenol			20	65-125	.5
2 - Methylphenol		20	65-125		.2
2 - Chlorophenol			20	65-125	.2
2 - Methyl 4,6 - Di	nitrophenol		20	65-125	.5
2 - Nitrophenol			20	65-125	.2
4 - Methylphenol		20	65-125		.5
4 - Chloro 3 - Meth	ylphenol		20	65-125	.2
4 - Nitrophenol			20	65-125	.5
Pentachlorophenol			20	65-125	1.0
Phenol			20	65-125	.2

Listed MDLs are from published EPA methods or based on historical TEG data.

^{1.} 2. Refer to Section 11.2 for definition of Precision.

Refer to Section 11.2 for definition of Accuracy, listed values from in-house data. Values also apply to surrogates.

TABLE 5-1 (continued)

187252

COMPONENT	MATRIX	METHOD	'PRECISION %RPD (Max)	ACCURACY %Recovery	MDL
	Groundwater	EPA 3510/608			
PCB'S	Surface water Effluents		·		ug/L
PCB 1016			20	65-125	20
PCB 1221			20 ·	65-125	50
PCB 1232			20	65-125	20
PCB 1242			20	65-125	15
PCB 1248			20	65-125	15
PCB 1254			20	65-125	10
PCB 1260			20	65-125	10
4,4 -DDD			20	65-125	0.05
4,4 -DDE			20	65-125	0.05
4,4 -DDT			20	65-125	0.05
Aldrin			20	65-125	0.05
alpha -BHC			20	65-125	0.05
beta -BHC			20	65-125	0.05
gamma -BHC			20	65-125	0.05
delta -BHC		•	20	65-125	0.05
Dieldrin		20	65-125		0.05
Endosulfan I			20	65-125	0.05
Endosulfan II			20	65-125	0.05
Endosulfan sulfate	1		20	65-1	0.05
Endrin			20	65-120	0.05
Endrin aldehyde			20	65-125	0.05
Heptachlor			20	65-125	0.05
Heptachlor epoxid	le		20	65-125	0.05
PCB'S	Soils and	EPA 3550/8080			mg/kg
	Sediments				
PCB 1016			20	65-125	300
PCB 1221			20	65-125	800
PCB 1232			20	65-125	300
PCB 1242			20	65-125	200
PCB 1248			20	65-125	200
PCB 1254			20	65-125	100
PCB 1260			20	65-125	100
4,4 -DDD			20	65-125	0.5
4,4 -DDE			20	65-125	0.5
4,4 -DDT			20	65-125	0.5
Aldrin			20	65-125	0.5
alpha -BHC			20	65-125	0.5
beta -BHC			20	65-125	0.5
gamma -BHC			20	65-125	0.5
delta -BHC			20	65-125	0.5
Dieldrin		20	65-125		0.5
Endosulfan I			20	65-125	0.5
Endosulfan II			20	65-125	0.5
Endosulfan sulfate			20	65-125	0.5
Endrin			20	65-125	0.5
Endrin aldehyde			20	65-125	0.5
Heptachlor	_		20	65-125	0.5
Heptachlor epoxid			20	65-125	0.5
	ublished EPA methods or be				

^{1.} Refer to Section 11.2 for definition of Precision.

^{2.} Refer to Section 11.2 for definition of Accuracy, listed values from in-house data. Values also apply to surrogates.

Section 6 Revision 1 Date 1/1/93 Page 1 of 2

6.0 SAMPLING PROCEDURES

6.1 SAMPLE CONTAINERS, PRESERVATION, HOLDING

Sample collection activities will not be performed by TEG. TEG prepares sampling packages consisting of appropriate containers with preservatives, coolers with ice, and chain-of-custody records. Sampling guidance documents are posted and available for reference at the fixed-base laboratory and mobile laboratories. Of key importance are procedures designed to reduce the potential for cross contamination of samples and volatilization of purgeables. A list of sampling containers and volumes, preservatives, and holding times is provided below in Table 6-1.

TABLE 6-1 CONTAINERS, PRESERVATION, AND HOLDING TIMES

PARAMETER	<u>MATRIX</u>	CONTAINER	PRESERVATION	HOLDING TIME
Organic Extractables	Soil/Waste Water	Glass Jar ⁱ 100mL	4°C 4°C	14 Days Until Extract.
		Glass Bottle		14 Days Until Extract. (30 Days extract to Analysis)
Purgeables (aromatics	Soil/Waste	Glass Jar ⁱ	4°C,	14 Days (ASAP)
and halocarbons)	Water	2x40inL VOA ²	4°C, HCl³ to pH < 2	14 Days (ASAP)

Approximately 250mL capacity jar with Teflon⁸-lined cap or drive-sample core tube with Teflon⁸-lined cap.

6.2 CLEANING

Most of TEG's glassware is of the disposable type and is not reused. For the small amount that is reused, the following cleaning procedures are used:

- wash with Alconox detergent and hot water:
- rinse thoroughly with tap water;
- rinse thoroughly with distilled water;
- rinse with reagent grade purge-and-trap methanol (retain rinsate);
- air dry;
- visually inspect; and
- analyze methanol rinsate for potential remnant contamination, as appropriate.

Volatile Organic Analysis Vials.

³ Reagent grade HCl is used for preservation.

Section 6 Revision 1 Date 1/1/93 Page 2 of 2

6.3 WASTE DISPOSAL

TEG is classified as a conditionally-exempt small quantity generator of hazardous wastes according to 40CFR Part 261. TEG generates less than 100 kg of hazardous waste per month. This hazardous waste is composed mostly of analytical solvents which are stored in acceptable containers and recycled by an approved facility.

Environmental samples analyzed by the laboratory are the property of the client, and in most cases involving on-site analysis, the samples are retained at the site. Samples which are analyzed at the laboratory are returned to the client with chain-of-custody report unless specific arrangements are made for disposal. Samples retained by TEG are disposed of by a licensed hazardous soil firm.

Section 7 Revision 1 Date 1/1/93 Page 1 of 11

7.0 SAMPLE CUSTODY PROCEDURES

7.1 SAMPLE CUSTODY OBJECTIVES

The purpose of this section is to provide an outline of sample custody protocol. This protocol is established for the purpose of providing traceability of samples and containers from point of origin to final disposition of the samples and to identify parties responsible for handling and transport of samples.

7.2 FIELD CUSTODY

TEG does not engage in field sampling services. Sampling is the responsibility of the consultant or client. The chain of custody protocol for the laboratory begins when sample containers are released by the laboratory for the sampler's use. If the client provides their own containers, TEG's chain-of-custody begins with sampling by the client. A TEG chain of custody record (Figure 7-1) is prepared by the TEG field chemist and provided to the sampler with the containers. A note regarding origin of sample containers is made under the "special instructions" heading on the form. The TEG chain of custody record is a three-color triplicate form. The bottom copy is retained by the sampler. The original (top) form and one copy are delivered to the laboratory with the samples. The completed original is then returned to the client with the analytical report. A copy of the completed original is maintained in the project file by the laboratory.

7.3 LABORATORY CUSTODY

<u>Sample Receipt</u>. Samples are received at TEG by the field chemist who is in charge of documentation for the job. Sample receipt and verification protocol is as follows:

- note presence, absence, and condition of the shipping container tape and custody seals:
- remove samples and documents from shipping vessel;
- organize samples by client ID numbers; and
- make notes regarding: temperature; breakage or opening; headspace in VOAs; septa orientation; labeling.

Sample Log-In. The following protocol is used for sample log-in:

- each batch of samples is assigned a project number;
- each sample is verified and matrix confirmed with the chain-of-custody record (Chain of Custody Record, Figure 7-1.);
- completeness of chain-of-custody report is noted;
- each sample is assigned a unique laboratory identification number;
- notes regarding condition received, integrity, seals, preservation, and any peculiarities are entered;

Section 7 Revision 1 Date 1/1/93 Page 2 of 11

- the Chain-of-Custody Report is signed, discrepancies noted, reasons for sample rejection noted;
- the samples are then stored in a limited access area and stored on ice or refrigerated, as appropriate; and
- samples are stored separately from standards, VOC samples are stored separately from other samples and standards.

The chemist logging in the samples, is responsible for noting irregularities and rejection of samples. Criteria for sample rejection include:

- elevated temperature, if cooling necessary;
- breakage or leakage;
- evidence of seal tampering;
- missing or duplicate samples;
- incomplete chain of custody record;
- missing labels;
- lack of preservation;
- insufficient sample for specified analyses;
- improper sample containers; and
- headspace in VOAs for VOC analysis.

<u>Sample Security</u>. Samples are stored in a limited access area of the mobile laboratory. Samples requiring cooling are stored in a refrigerator.

<u>Sample Distribution and Tracking</u>. Samples are signed out from storage by the field chemist or analyst. An Extraction Log (Figure 7-2) and an Analysis Log (Figure 7-3) are maintained by the analyst. On completion of analysis, or taking an aliquot from the sample, the remaining sample, and any excess extract or digest, is returned to storage.

Each form used in tracking and handling the sample requires time, date and initials or signature of persons performing the work. In the TEG mobile laboratory setting, the same field chemist is generally in charge of all phases of sample tracking from receipt to analysis. Although most samples are analyzed within a few hours of receipt, it is the responsibility of the field chemist to meet holding time constraints in the extraction and analysis of samples. Laboratory analytical reports include dates and time of sampling, extraction, and analyses subject to review by the QA Manager to assure that timing criteria have been achieved.

Samples and extracts are retained for 15 days following reporting, at which point they are returned to the client, via chain-of-custody record, or disposed of in an approved manner consistent with the nature of the sample. Longer storage may occur by prior arrangement with the client. Sample disposition is conducted in accordance with waste disposal guidelines described in Section 6.3.

Section 7 Revision 1 Date 1/1/93 Page 3 of 11

Samples transferred to other laboratories are packaged at appropriate temperatures and shipped with chain-of-custody record. The same information required in the above-listed protocol for sample custody applies to samples and extracts shipped to other laboratories.

<u>Document Control</u>. TEG maintains a document control program to provide guidelines in reducing the potential for document loss and mix-ups. The data generator (field chemist) is responsible for initiating a file for each batch of samples on delivery to the laboratory. The following procedures are implemented as part of the document control process:

- a job file is labeled with job number and client name;
- a Project / Report Checklist Form (Figure 7-4) is initialized in duplicate;
- on completion of each task, or at the end of each day, all paperwork and forms generated for the project are transferred to the master file with the contents noted on the inventory form;
- all magnetic computer media is to be backed up to a floppy disk which is stored with the job file; and
- on completion of the project, the report preparer will close the file by verifying all contents are present and sending the file on for signature.

7.4 ELECTRONIC DATA RECORDS

TEG uses microcomputers for data storage and reporting purposes. Initial chromatographic data gathering is performed by integrators which print a chromatogram and report integration. Parameters of integration are established for each analysis. It is the responsibility of the field chemist to evaluate each chromatogram with regard to correct integration, peak identification, and calibration. Verified numeric data from the chromatogram is then manually entered into a spreadsheet on the microcomputer. A spreadsheet file is created from a generic starter file for each project and analytical method. The spreadsheet data is saved on a floppy disk which is stored in the job file.

Section 7
Revision 1
Date 1/1/93
Page 4 of 11

Figure 7-1. Chain of Custody Record

Ý.	THANSGLOOM
21	ENVIRONMENTAL
tea	GEOGIEMISTAY, INC.
· J	

CHAIN-OF-CUSTODY RECORD

ſ						_						_			Г																	
CLIENT:						_				_		_		—												PAGE			OF_		<u> </u>	_
												_	_	_	TI	EG	PR	DJE	СТ	•:	_			_			—					_
PHONE:				FAX:		_				_		—		_	L	oc.	ATIC	ON:	_			_		_			_					_
CLIENT PROJE	CT # :_			PROJECT M								_		_	c	סנג	ECT	OR	: <u> </u>	_	_						_	COL	LECTIO			_
Sample Number	Depih	Time	Sample Type	Container Type	SA.	, o /															\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\					// FEE	ON C.	TES.		Total Number	O Container	Hote Bumber
	1		1		1	ĺ	Ĺ	Ĺ			Ĺ	Ĺ						أ					ĺ	Í	ĺ					1	Ĭ	
							Ī				Ī	匚	匚				\Box	Ì			i			Ţ	j					T	\Box	
						Ī			\Box				<u>L</u>											1	ĺ						1	
					┖	L	<u> </u>	<u> </u>	Ц		_	L	Ļ	1 1			_	_				<u> </u>	<u> </u>	Ļ	1					ᆚ_	1	
					<u> </u>	┖	_	L			L	L	上	11	_	_	_	_			L	<u> </u>	L	Ļ	┖					1	_	
					┞-	Ļ.	1	╙		_	L	╙	ᆜ	1	_	_	_	_	_	Ш	L	_	<u> </u>	Ļ	<u> </u>					+		_
	<u> </u>				<u> </u>	<u> </u>	<u> </u>	<u> </u>		_	Ļ.	⊢	╀	\sqcup	_	_ļ	4	4	4		_	<u> </u>	<u> </u>	Ļ	<u> </u>					+	_	
				•	┞	┞	!			_	L	Ë	!		4	_ !	4	4	긕		_		Ļ	Ļ	Ļ					+-	+	
					┼	├	<u> </u>			-	_	├-	⊢	H	+	<u> </u>	+	4	-		_		┝	1	÷	_				+-	-	!
					⊢	╁	+-	Н	-	\dashv		┢	⊢	1	+	ᅥ	ᆉ	╣	닉	-	H	\vdash	╀	╀	1		—	_		+	$\frac{1}{1}$	-
					╁	\vdash	+-	Н	-	-	_	┢	┝	H	+	┥	+	+	┥		H	_	╁	╀	╁					+	÷	-
	- +				\vdash	╁	H	\vdash	\dashv	⊣	Н	Н	H	H	+	┪	\dashv	7	\dashv		_	Н	H	+	t		_			+-	7	
		$\neg \neg$			\vdash	İ	İ	П	-	┪		Г	一	H	\dashv	寸	Ť	7	\dashv			1	H	Ϋ́	<u>†</u>					Ť		
			1		\vdash	İ	匸	П		\exists		П	厂	H	\dashv	i	i	寸	7			İ		T	Ì		_			\dagger	二	_
						Π		П	\Box	٦			Г		T		Î	- 1	\neg					Ī	1					T	1	_
					<u> </u>	<u> </u>	i						\Box		\perp		\Box								Ī				_			
					<u> </u>	_	1		ı	۷		Ш	L.	\sqcup				_					<u> </u>	L	1					丄	1	
RELINOUISHED BY.	(5-gnatur	e)	DATEA	ME RECEIVED) BY	(54	~12	46)		DAT	I E/T	IME		<u> </u>	_		54	_	_		_	_		4	4	LABORAT	ORY	NOTE	ES:			- 1
RELINOUISHED BY	(S-gnatur	e)	DATEZIA	ME AECEIVE	BY.	(\$4	;^a~	ee]		DA	TEA	i indi		<u>Ch</u>	AIN	01	CU	\$ 70	DY	SE	AL:			-	Ⅎ							
	s	AMPLI	EDISPO	SAL INSTRUCTI	ON	s		_		-	-	_	_	_			GC		_	_		011	_	+	ᅱ							
				en C Between	_	_		_	_			_			<u> </u>	V E 1	, 00	,,,,	<u> </u>	J 14 L	<i>7.</i> / _	ΨĻ	_	4	-1							

Section 7

Revision 1

Date 1/1/93

Page 5 of 11

Figure 7-2. Extraction Log

	t			*		TEG	EXTRACTION OF THE PROPERTY OF	ON LCG			
CLIENT	:							CLI	ENT PRO	JECT #: _	
DATE:				_	TEG	≠ :			ANA	LYST/S: _	
LOCATIO) 				<u>:</u>				_		
EPA Met 8010/601 3020/602 8015 418.1 Cther	{	Wa	ter	rix:		H H H	Solvent: ethylene arbon S2 exame ethamol reon-113	c1 _	C S S	Method: and Shake ont Lig/L chication oxhlet ther	iq
SAMPLE #	TIH	E		ESTI NCEN		D YON		SPIKE		INITIAL	SCLVENT
						Unk		Con	Vol	ç::s	7.15
		-				<u> </u>		<u> </u>			
<u> </u>		_				<u> </u>		<u> </u>			
					<u> </u>	<u> </u>		<u> </u>			<u></u>
	<u> </u>	<u> </u>			<u> </u>			<u> </u>	<u> </u>		
		\perp			ļ 				<u> </u>		
								<u> </u>	<u> </u>		
		_						<u> </u>	1		
	 	1				<u> </u>					
	<u> </u>	+	_					<u> </u>	<u> </u>		
		_	[[]			<u> </u>	<u> </u>	
		1	_					<u>†</u>		- !	
	<u> </u>	4	_							<u></u>	
	<u> </u>	4	_								
		+									
	 		_							1	
	ļ	+	_				_				
		\bot	_								
<u> </u>		\bot	_								
		\bot	_								
 -		-		-	.						

Section 7 Revision 1 Date 1/1/93 Page 6 of 11

Figure 7-3. Analysis Logs

teg

TRANSGLOBAL

ENVIRONMENTAL .

GEOCHEMISTRY, INC.

wy				a cooleman,
	TEG J08#:	CLIENT:	CLIENT JC	
	DATE:	ANALYST:	instrum:	 .
SAMPLE:				
MATRIX:	-			
RUN #:	i !			
ANALYTE: TIME:				1 1
1.1 DICHLOROETHENE	i			1
METHYLENE CHLORIDE				
,2 iransDiCHLOROETHENE I	1	i I	1 1	
1 DICHLOROETHANE		i i	1	
ICHLOROFORM I	! 1		1	
1.2 cisDICHLOROETHENE	T	i i	: 1	
,1.1 TRICHLOROETHANE I	1			1 1
ARBON TETRACHLORIDE			1 1	
2 DICHLOROETHANE	1	1		
TRICHLOROETHENE				
1.2 DICHLOROPROPANE	1 1			
ROMODICHLOROMETHA I		i_ i	i i	
ICHLOROPROPENE	1 1			
.1.2 TRICHLOROETHANE		1	;	
TETRACHLOROETHENE 1	1 _1 _		1 1	
TETRACHLOROETHANE I	1 1			
HLOROBENZENE	ii	T I	ii	
ENZENE	1 1	i i		
OLUENE			11	
ETHYLBENZENE			i ī	
M&P XYLENE		1	1	
XYLENE			<u> </u>	
OTAL XYLENES		Ī	· _ 1	

EPA METHOD 601/S010 AND 602/S020 ANALYSES LOG

Section 7 187261 Revision 1

Date 1/1/93
Page 7 of 11

Figure 7-3. Analysis Logs Continued

TRANSGLOBAL

'ceg

EPA METHOD 60\$ 80\$0 AND PCB ANALYSES LOG

ENVIRONMENTAL GEOCHEMISTRY, INC.

	TEG JOB#:	CLIENT:	CLIENT JOB#		
	DATE:	AN ST:	INSTRUM:		
SAMPLE:			-		
MATRIX:				11	
RUN #: 1		<u> </u>	1	<u> </u>	
ALYTE: TIME:	<u> </u>	<u> </u>			
ha-BHC t I				1	
nma-BHC			1	1	
ACHLOR	<u> </u>				
. <u>1N</u>		<u>l</u>	! !	<u> </u>	
HC a-BHC	<u>i</u>	1	i :	i	i
a-BHC	į l	1	t	- 1	
PTACHLOR EPONIDE	<u></u>	1 1	1	1	
SULFANI		i	1		
DE 1 (i l	1	1	1	
DRIN	<u> </u>		1 1	1	
DRIN	i		1	1	1
DDD	1	1	7 ;	1	
SULFAN II	4 1	1	-	I	
TC:	1	; 1	!	1	1
INALDEHYDE	1		1		
DOSULFAN SULFATE	<u> </u>	i 1			1
3UTYL CHLORENDATE	- i	1 1	i	- i	
YPE IDENTIFIED:	1 1	1 i			1
YPE IDENTIFIED:				1	i —

Section 7 187262 Revision 1 Date 1/1/93 Page 8 of 11

Figure 7-3. Analysis Logs Continued

of_					YSIS LCG	TREH
CLIENT: DATE:						ENT PROJECT #:
CONDITI						INSTRUMENT: BUCK IR
						<
SAMPLE #	TIME	_ RUN	ABS	DIL	conc	COMENTS
	<u> </u>	1 1		<u> </u>		1
	+	<u> </u>				
	+		_			
	† <u>-</u>					
			_	_		
					•	
	<u> </u>					
	<u> </u>	<u> </u>			<u> </u> 	
	1		_			
<u> </u>	-					
	1 1					

Section 7 187260 Revision 1 Date 1/1/93 Page 9 of 11

Figure 7-3. Analysis Logs Continued

01				ΞPλ	8015	i		
CLIENT	:					CLIENT PRO	JECT #:	
DATE:			_ 15	G ≠:		ANA	LYST/S:	
LCCATI	ON:							
CONDIT	IONS:					STRUMENT:	SHIMADZU	GC14A
Gas	Factor:				Diesel	Factor:		
SAMPLE #	TIME	ಸರಾಗ	VOL	GAS AREA	GAS CONC	DIESEL AREA	CONC	COMMENTS
					<u> -</u>			·
					<u> </u>			
					<u> </u>			
						<u> </u>		
							<u> </u>	<u> </u>
					1	<u> </u>	<u> </u>	
					<u> </u>	<u> </u>	ļ	
· 					1	1	ļ	<u> </u>
	1 1						<u> </u>	
						<u> </u>	<u> </u>	
			‡		ļ	<u> </u>	<u> </u>	
							-	
					<u> </u>			<u> </u>
					<u> </u>		1	
							 	
								
							1	
							1	
	1 [

Section 7 Revision 1 Date 1/1/93 Page 10 of 11

Figure 7-3. Analysis Logs Continued

of				TEG	ANAL EPA	YSIS LO	:G			STEX
CLIENT	:					·	CLIE	T PRO	JECT #:	
LOCATIO								•		
CONDITI							NSTRUM	ENT:	SHIMADZ	U GCSA
SAMPLE /	Time	Run	!		Tol		Eth	MSP Xyl		Comments
					<u> </u>					
	<u> </u>					<u> </u>		$oxed{oxed}$		
	1 1									
									i	
					1					
							T			
			1			İ			 i	
				1					- i	
		1					$\neg \dagger$		 	
		十	一	- +	- 		_			
		$\neg \uparrow$	\dashv	 	<u>'</u> -		-+			
			_	T	- 		-+			
				- 	$\neg +$		$\neg +$		- +	
							-+		- -	

Section 7 187265 Revision 1 Date 1/1/93 Page 11 of 11

Figure 7-4. Project / Report Checklist Form

	_	
		PROJECT / REPORT CHECKLIST
NEED	DONE	
		Preliminary Data
1 -	_	☐ To Client on-site ☐ Faxed to client
		TEG additional analyses
-		□8010 □8015 □8020 □418.1 □ Cther
		Outside analyses:
1 -	-	DATI DIQAL Dither lab
1 a	Q	Spreadsheet transferred to current (loppy cisk
anaaaa	ä	Data tables completed
1 5	ä	O LOC Mately Called & Mathed Stants data technical
1 5	ä	QA/QC-Matrix Spike & Method Blank data Included Cover letter
1 2		Centilication number - correct/match
1 2	ä	
1 4 .	u	
1 0.		□ Client □ Other party □ Accounting □ File-copy
	<u> </u>	
0	0	Final Review
"	0	Report to citent
	_	□ Mail □ Deliver
"	۵	Additional Report(s) to:
	_	□ DOHS □ HMMD □ Orange Co. DHS □ Cifer
		Report/Data to be added/included with:
ì		TEG Project #
		Client/Company

8.0

ANALYTICAL PROCEDURES

187266

8.1 METHOD REFERENCES

The analytical methods used at TEG are based on procedures in the following references:

- U.S. EPA Document SW-846, Test Methods for Evaluating Solid Waste, Third Edition, November, 1986, and Draft Revision 1, December 1987;
- U.S. EPA Document 600/4-79-020, Methods for Chemical Analysis of Water and Wastes, March 1983;
- U.S. Code of Federal Regulations 40, Protection of Environment, Part 136, Guidelines Establishing Test Procedures for the Analysis of Pollutants, July 1, 1989;
- U.S. Code of Federal Regulations 40, Protection of Environment, Part 141, National Primary Drinking Water Regulations, July 1, 1989; and
- California State Water Resources Control Board, Division of Water Quality, Leaking Underground Fuel Tank Field Manual, December 17, 1987.

This section of the document provides an outline of the analytical methods referenced in Table 5-1. Data packages generated by TEG for certification of EPA methods are available upon request.

8.2 EPA METHODS 601 and 8010

<u>Introduction</u>. The purpose of this procedure is to describe the GC method for determination of purgeable halogenated hydrocarbons in soils, solid wastes, and waters using the Hall Detector. Details of the method are found in EPA Document SW-846. Presented below is an overview of the method in terms of calibration, operating conditions, compound identification, and calculations. This method can be modified to include analysis of EDB with an electron capture detector in accordance with the requirements of certain states.

Calibration. Calibration may be accomplished by either internal or external standard techniques. Standards are prepared at five concentrations spanning the linear range of the instrument. The standards are then analyzed and the response factor evaluated for linearity. If linearity is not achieved a second set of analyses is run. If linearity is still not achieved, a second set of standards is prepared and the process repeated. Linearity is acceptable when response factor values are within 15% standard deviation for internal standard calculations or if the mean residual from the linear regression is within 15%.

<u>Water Sample Preparation</u>. For aqueous samples, a 5mL aliquot is generated and spiked with internal standards and (or) surrogates in a gas tight syringe. Purge and Trap operation is performed in general accordance with EPA method 5030. The sample is purged for eight minutes at a flow rate of 30mL per minute.

Section 8 Revision 1 Date 1/1/93 Page 2 of 16

<u>Soils and Solid Wastes</u>. A solvent extraction is first performed using methanol. The methanolic extraction is achieved by placing ten grams of the soil sample into a 40mL VOA (volatile organic analysis) vial, adding 10uL of surrogate soil spike (1000ng/uL in methanol), and 10mL of reagent purge-and-trap grade methanol. The VOA is then hand shaken for two minutes and placed in a water-bath sonicator for ten minutes. Depending on anticipated concentration, 10 to 50uL of the extract is then added to 5mL of volatile free water in the purge and trap or 3ul of the extract is directly injected into the gas chromatograph.

Purge and Trap Conditions.

Instrument:

Tekmar LSC 2000

Purge flow:

20mL/min to 30mL/min

Purge time:
Dry purge time:

8 minutes

Desorb time:

8 minutes 3 minutes

Desorb temperature:

275 degrees C

Bake time:

10 minutes

Bake temperature:

290 degrees C

Gas Chromatography.

Instrument:

Gas Chromatograph

Column:

50 to 100 meter by 0.053mm, RT 502.2, megabore

capillary.

Carrier flow:

Helium at 15mL/min.

Detector:

Electrolytic Conductivity detector.

Detector temperature:

225 degrees C

Injector temperatures:

175 degrees C

Column oven:

45 degrees C for 2 minutes

45 to 200 degrees C at 5/minute

<u>Standard Preparations</u>. Primary 8010 standards at 100mg/L (ppm)in methanol are purchased from a certified supplier.

Secondary Standards (lng/uL): dilute primary standard by 100 times (100uL to 10mL).

Water Matrix Spikes. For lug/L (1 ppb) water concentration, add 5uL of lng/uL (ppm) secondary 601 standard to 5mL of water (1,000 times dilution).

Soil Matrix Spikes. For 1mg/kg (1 ppm) soil concentration, add 100uL of 100 ng/uL primary standard to ten grams of soil.

Typically, the following 10 compounds are quantified on matrix spikes covering early, middle, and late eluters: methylene chloride, 1,2-dichloroethane, 1,1-dichloroethane, chloroform, 1,1,1-trichloroethane, carbon tetrachloride, trichloroethylene, 1,1,2-trichloroethane, tetrachloroethylene, and tetrachloroethane. Other compounds are used depending upon the target compounds executed.

Section 8
Revision 1
Date 1/1/93
Page 3 of 16

Calculations of Sample Concentrations.

Water Samples

Concentration (ug/L) = [(Ax * A)/(As * Vs)] * D

Soil Samples

Concentration (ng/g) = [(Ax * A * Vt * D)/(As * Vi * W)]. Where:

Ax is the area counts for the sample - method blank

A is the amount of standard injected (ng)

As is area counts for the standard

Vi is the volume of extract injected (uL)

D is the dilution factor on the sample

Vt is the volume of total extract (uL)

Vs is the volume of water extracted (mL)

W is the weight of soil extracted (g)

For the standard procedures employed by TEG, these equations reduce to:

Water Samples: Conc (ug/L; ppb) = measured mass (ng)/5

<u>Compound Identification</u>. This method is used to identify and quantify the analytes listed in Table 5-1.

8.3 EPA METHODS 602 and 8020

Introduction. The purpose of this procedure is to describe the GC method for determination of volatile aromatic hydrocarbons in soils, solid wastes, and waters using a photoionization detector. Details of the method are found in EPA Document SW-846. Presented below is an overview of the method in terms of calibration, operating conditions, compound identification, and calculations. This method can be modified to include Methyl tert-Butyl Ether (MTBE) in accordance with the requirements of certain states.

<u>Calibration</u>. Calibration may be accomplished by either internal or external standard techniques. Standards are prepared at five concentrations spanning the linear range of the instrument. The standards are then analyzed and the response factor evaluated for linearity. If linearity is not achieved a second set of analyses is run. If linearity is still not achieved, a second set of standards is prepared and the process repeated. Linearity is acceptable when response factor values are within 15% standard deviation for internal standard calculations or if the mean residual from the linear regression is within 15%.

Section 8
Revision 1
Date 1/1/93
Page 4 of 16
187269

Sample Preparation

<u>Water Samples</u> A 5mL aliquot of water is spiked with 5uL of 1 ng/uL surrogate standard (chlorobenzene) and extracted by purge and trap.

<u>Soil Samples</u> 10 grams of soil are weighed to the nearest 0.1 gram and placed into a clean VOA vial with 10mL of purge and trap grade methanol or freon-113. The VOA vial is shaken for 2 minutes, placed into a water bath and sonicated for 10 minutes. After sonication, the slurry is allowed to settle and 10 to 50uL of solvent are withdrawn, added to 5mL of water, and extracted by purge and trap or 3ul of the extract is directly injected into the gas chromatograph.

Purge and Trap Conditions.

Instrument:

Tekmar LSC 2000

Purge flow:

20mL/min to 30mL/min

Purge time:

8 minutes

Dry purge time:
Desorb time:

8 minutes 3 minutes

Desorb temperature:

275 degrees C

Bake time:

10 minutes

Bake temperature:

290 degrees C

Gas Chromatography.

Instrument:

Gas Chromatograph

Column:

30 meter by 0.053mm, DB-1 or DB-5, megabore

capillary.

Carrier flow:

Helium at 15mL/min.

Detector:

Photoionization detector.

Detector temperature:

225 degrees C

Injector temperatures:

175 degrees C

Column oven:

45 degrees C for 4 minutes

45 to 175 degrees C at 10/minute

Standard Preparations. Primary 8020 standards at 100mg/L (ppm)in methanol are purchased from a certified supplier.

Secondary Standards (lng/uL): dilute primary standard by 100 times (100uL to 10mL).

Water Matrix Spikes. For lug/L (1 ppb) water concentration, add 5uL of lng/uL (ppm) secondary 602 standard to 5mL of water (1,000 times dilution).

Soil Matrix Spikes. For lmg/kg (1 ppm) soil concentration, add 100uL of 100 ng/uL primary standard to ten grams of soil.

Matrix spikes consist of the following compounds: benzene, toluene, chlorobenzene, ethylbenzene, m-xylene, o-xylene, and p-xylene.

Section 8 Revision 1 Date 1/1/93 Page 5 of 16

Calculations of Sample Concentrations. Water Samples

Concentration (ug/L) = [(Ax * A)/(As * Vs)] * D

Soil Samples

Concentration (ng/g) = [(Ax * A * Vt * D)/(As * Vi * W)]. Where:

Ax is the area counts for the sample - method blank

A is the amount of standard injected (ng)

As is area counts for the standard

Vi is the volume of extract injected (uL)

D is the dilution factor on the sample

Vt is the volume of total extract (uL)

Vs is the volume of water extracted (mL)

W is the weight of soil extracted (g)

For the standard procedures employed by TEG, these equations reduce to:

Water Samples: Conc (ug/L; ppb) = measured mass (ng)/5

Soil Samples: Conc (mg/kg; ppm) = measured mass (ng)/Vi

<u>Compound Identification</u>. This method is used to identify and quantify the analytes listed in Table 5-1.

8.4 EPA METHOD 8015 TPH

Introduction. The purpose of this procedure is to describe the GC method for determination of Total Petroleum Hydrocarbons (TPH) in soils, solid wastes, and waters using a flame ionization detector. Details of the method are found in EPA Document SW-846 and the California LUFT Manual. The method is applicable to determination of gasoline, diesel, and other volatile and semi-volatile fuels in water, soils, and wastes. This method is approved by California Department of Health Services as a quantitative method for the determination of total fuels content.

<u>Calibration</u>. Calibration is accomplished by using external standard techniques. Standards are prepared at three concentrations spanning the linear range of the instrument. Standards composed of the anticipated fuel product to be analyzed are used for the calibration. The standards are then analyzed and the response factor evaluated for linearity. If linearity is not achieved a second set of analyses is run. If linearity is still not achieved, a second set of standards is prepared and the process repeated. Linearity is acceptable when response factor values are within 15% standard deviation for internal standard calculations or if the mean residual from the linear regression is within 15%.

Section 8 Revision 1 Date 1/1/93 Page 6 of 16

Sample Preparation and Extraction

<u>Water Samples</u>. A 33mL aliquot of water is extracted with 1mL of solvent by shaking vigorously for 5 minutes in a clean 40mL VOA vial. The mixture is allowed to settle and 3uL of solvent are withdrawn and injected into the gas chromatograph.

<u>Soil Samples</u>. 10 grams of soil are weighed to the nearest 0.1 gram and placed into a clean VOA vial with 10mL of solvent and NaSO4. The VOA vial is shaken for 2 minutes, and sonicated for 10 minutes. After sonification, the slurry is allowed to settle and 3uL of solvent are withdrawn and injected into the gas chromatograph.

Gas Chromatography.

Instrument:

Gas Chromatograph with FID

Column:

30 meter by 0.053, DB-1, megabore capillary.

Carrier flow:

Helium at 15mL/min.

Injector temperature:

175 degrees C

Detector temperatures:

225 degrees C 45 degrees C for 4 minutes,

Column oven:

45 to 205 degrees C at 10/min,

Hold at 205 degrees C for 10 minutes.

Standard Preparation.

Substock Calibration Standards (in trichlorotrifluoroethane)

Low calibration standard (20 ng/uL): add 0.25uL gasoline to 10mL solvent. Mid calibration standard (200 ng/uL): add 2.5uL gasoline to 10ml solvent. High calibration standard (2000 ng/uL): add 25uL gasoline to 10mL solvent.

Low calibration standard (50 ng/uL): add 0.55uL diesel to 10mL solvent. Mid calibration standard (500 ng/uL): add 5.5uL diesel to 10ml solvent. High calibration standard (5000 ng/uL): add 55uL diesel to 10mL solvent.

Soil Matrix Spikes

Gasoline: add 2.5uL pure gasoline to 10 grams soil = 200mg/kg (ppm) Diesel: add 5.5uL pure diesel to 10 grams soil = 500mg/kg (ppm)

Water Matrix Spikes. For 2mg/L (1 ppm) water concentration, dilute substock standards by 100 times.

Calibrations of Sample Concentrations

Water Samples

Conc (ug/L) = (Ax * A * Vt * D)/(As * Vi * Vs)

Section 8 Revision 1 Date 1/1/93 Page 7 of 16

Soil Samples

Conc (ng/g) = (Ax * A * Vt * D)/(As * Vi * W)

Where:

Ax is the area counts for the sample - method blank

A is the amount of standard injected (ng)

As is area counts for the standard

Vi is the volume of extract injected (uL)

D is the dilution factor on the sample

Vt is the volume of total extract (uL)

Vs is the volume of water extracted (mL)

W is the weight of soil extracted (g)

8.5 EPA METHOD 418.1

Introduction. The purpose of this procedure is to describe the method for determination of Total Recoverable Petroleum Hydrocarbons (TRPH) in soils, solid wastes, and waters using an infrared spectrometer. Details of the method are found in EPA Document SW-846. The method is applicable to determination of petroleum hydrocarbons in water, soils, and wastes. This method is approved by EPA as a quantitative method for the determination of total fuels content.

Calibration. Calibration for the method is accomplished by using external standard techniques. Standards are prepared at three concentrations spanning the linear range of the instrument. Standards composed of the anticipated fuel product to be analyzed are used for the calibration. The standards are then analyzed and the response factor evaluated for linearity. If linearity is not achieved a second set of analyses is run. If linearity is still not achieved, a second set of standards is prepared and the process repeated. Linearity is acceptable when response factor values are within 15% standard deviation for internal standard calculations or if the mean residual from the linear regression is within 15%.

Sample Preparation and Extraction

<u>Water Samples</u>. A 33mL aliquot of water is extracted with 3mL of solvent by shaking vigorously for 5 minutes in a clean 40mL VOA vial. The mixture is allowed to settle and 2ml of solvent are withdrawn and analyzed on the Infrared Spectrometer.

<u>Soil Samples.</u> 10 grams of soil are weighed to the nearest 0.1 gram and placed into a clean VOA vial with 10mL of solvent and NaSO4. The VOA vial is shaken for 2 minutes, and sonicated for 10 minutes. After sonification, the slurry is allowed to settle and 2ml of solvent are withdrawn and analyzed on the Infrared Spectrometer.

Section 8 Revision 1 Date 1/1/93 Page 8 of 16

Equipment.

Instrument:

Buck Scientific HC-404

Standard Preparation

Substock Calibration Standards (in trichlorotrifluoroethane)

Low calibration standard (50ng/uL): add .55uL diesel or oil to 10mL solvent. Mid calibration standard (500ng/uL): add 5.5uL diesel or oil to 10ml solvent. High calibration standard (5000ng/uL): add 55uL diesel or oil to 10mL solvent.

Soil Matrix Spikes Add 5.5uL pure diesel or oil to 10 grams soil = 500mg/kg (ppm)

Water Matrix Spikes For 5mg/L (5 ppm) water concentration, dilute 500 ppm substock standards by 100 times.

Calculations of Sample Concentrations

Water Samples

Conc
$$(ug/L) = (Ax * A * Vt * D)/(As * Vi * Vs)$$

Soil Samples

Conc
$$(ng/g) = (Ax * A * Vt * D)/(As * Vi * W)$$

Where:

Ax is the area counts for the sample - method blank

A is the amount of standard injected (ng)

As is area counts for the standard

Vi is the volume of extract injected (uL)

D is the dilution factor on the sample

Vt is the volume of total extract (uL)

Vs is the volume of water extracted (mL)

W is the weight of soil extracted (g)

8.6 EPA METHOD 610/8100

Introduction. This method can be used in determining the concentration of PAH's present in soil and water samples. This procedure involves extracting the PAH's from a 10 gram sample of soil or a 30 gram sample of water using freon. The extract is then analyzed by gas chromatography using an flame ionization detector. The detection limits for each PAH using this procedure are listed in Table 5-1.

Calibration. Calibration may be accomplished by either internal or external standard techniques. Standards are prepared at three or five concentrations spanning the linear range of the instrument. The standards are then analyzed and the response factor evaluated for linearity. If linearity is not achieved a second set of analyses is run. If linearity is still not achieved, a second set of standards is prepared and the process repeated. Linearity is acceptable when response factor values are within 15% standard deviation for internal standard calculations or if the mean residual from the linear regression is within 15%.

Apparatus and Reagents

- Freon gas chromatographic grade
- Sodium sulfate anhydrous
- 10 N Sodium hydroxide solution
- PAH stock solutions 2000 ppm in methylene chloride.
- Surrogate spiking solution 1000 ppm of 2-Fluorobiphenyl in 2-propanol.
- Silica gel cartridges prepared with activated 100/200 mesh silica gel.
- VOA vials 40mL glass w/polyethylene screw caps.
- Shimadzu 14A GC with an flame ionization detector using a PTE-5 QTM, 15 moder, 0.53 mm diameter column, 0.5 μm film and Helium as carrier gas.

Injection port temp - 250 degrees C

Detector temp - 300 degrees C

Column temp - 75 degrees C/2 min. to 300 degrees C at 12/min 3 µl injection for analysis

Sample Preparation. Tare a 37.5ml VOA vial on a top loading balance and transfer 10 gram +/-.1 gram of soil sample in the vial. Add enough anhydrous sodium sulfate to vial until the soil becomes free flowing. Spike the soil sample by adding 10 μ l of the surrogate spiking solution directly into the soil. Add 10ml of freon and cap vial. Shake sample well for 2 minutes and then sonicate for 10 minutes. Sample can now be analyzed by direct injection into the gas chromatograph.

For water samples, place 30.0 grams of water to be analyzed in a voa vial. Add enough base to water to bring pH to above 11. Spike the water sample by adding 3 μ l of the surrogate spiking solution directly into the water. Transfer 2mls of freon to vial, cap and shake vigorously for 5 minutes. The freon layer (bottom) can now be analyzed directly. Results must be divided by 15 due to the concentration factor of the extraction procedure.

If any extract requires cleanup procedure, transfer 1-2ml of the extract through a silica gel column that has been prewashed with 2-3ml of freon. Discard the first 0.5mL of the extract that elutes from the cartridge, then collect the remaining fraction in a VOA vial. The extract is now ready for analysis by direct injection into the gas chromatograph.

Section 8 Revision 1 Date 1/1/93 Page 10 of 16

<u>OA/OC</u> Analyses. At the beginning of every 12 hour shift a continuing calibration is to be run to verify that the calibration is still valid. The recommended concentration of this standard is 1 ppm of each component. The determined concentration of each PAH must be within 80% to 120% for the calibration to be considered valid. If these limits are not met a 4 point calibration is to be performed to reestablish calibration. The standard concentrations that are recommended are 0.5, 2.0, 5.0 and 10.0 ppm for both water and soil analysis.

For each set of sample analyses, a method blank is to be analyzed. A method blank consists of the anhydrous sodium sulfate shaken with 10ml of freon in a VOA vial. If sample cleanup was performed on any of the samples analyzed, a method blank must be analyzed in which the method blank was passed through a prepared silica gel column.

After each 10 samples are analyzed, a PAH matrix spike and matrix spike duplicate are to be extracted and analyzed. The recommended concentration to spike soil samples is 1 ppm. This is performed by spiking 10 + / - .1 gram of a soil sample with 15 μ l of the 2000 ppm PAH stock solution. The recommended concentration to spike water samples is .2 ppm. This is performed by spiking 30 + / - .1 gram of a water sample with 3 μ l of the 2000 ppm PAH stock solution. The spiked sample is then extracted according to the procedure outlined above. After analysis of the matrix spike and matrix spike duplicate, the percent recovery is to be determined as followed:

The acceptable % recovery limits are 65% to 125%. In addition to determining the percent recovery, the % RPD is to be determined and reported. To calculate the % RPD use the following calculation:

$$\% RPD = \% Rec_{MS} - \% Rec_{MSD}$$

$$\frac{(\% Rec_{MS} + \% Rec_{MSD})/2)}{(\% Rec_{MS} + \% Rec_{MSD})/2}$$

This value is to be reported along with the % Recovery of the matrix spike and matrix spike duplicate.

8.7 EPA METHOD 608/8080

Introduction. This method can be used in determining the concentration of pesticides and PCB's present in soil and water samples. This procedure involves extracting the pesticides and PCB's from a 10 gram sample of soil or a 30 gram sample of water using hexane. The extract is then analyzed by gas chromatography using an electron capture detector. The detection limits for each pesticide and PCB using this procedure are listed below.

Section 8
Revision 1
Date 1/1/93
Page 11 of 16

<u>Calibration</u>. Calibration may be accomplished by either internal or external standard techniques. Standards are prepared at three or five concentrations spanning the linear range of the instrument. The standards are then analyzed and the response factor evaluated for linearity. If linearity is not achieved a second set of analyses is run. If linearity is still not achieved, a second set of standards is prepared and the process repeated. Linearity is acceptable when response factor values are within 15% standard deviation for internal standard calculations or if the mean residual from the linear regression is within 15%.

Apparatus and Reagents.

- Hexane pesticide grade
- Sodium sulfate anhydrous
- PCB stock solutions 200 ppm in methanol
- Pesticide stock solution 20 ppm and 2 ppm in methanol
- Surrogate spiking solution 100 ppm Dibutylchlorendate in methanol.
- Florisil cartridges Supelco LC-SI 6ml tubes
- VOA vials 40mL glass w/polyethylene screw caps.
- Shimadzu 14A GC with an electron capture detector using a Rtx-5, 30 meter, 0.53 mm diameter column with 1.5 μ m coating thickness. Confirmation column for pesticides is a DB-17, 30 meter, 0.53 mm diameter column with 0.5 μ m coating thickness.

Injection port temp - 300 degrees C

Detector temp - 300 degrees C

Column temp - 150 degrees C /3 min to 290 degrees C at 4/min 3 µl injection for analysis

Nitrogen for carrier gas

Sample Preparation. Tare a 40mL VOA vial on a top loading balance and transfer 10 gram +/-.1 gram of soil sample in the vial. Add enough anhydrous sodium sulfate to vial until the soil becomes free flowing. Spike soil with $10 \mu l$ of the surrogate spiking solution. Add 10ml of hexane and cap vial. Shake sample well for 2 minutes and then sonicate for 10 minutes. Sample can now analyzed by direct injection into the gas chromatograph.

For water samples, place 30.0 grams of water to be analyzed in a voa vial. Spike sample with 3 μ l of surrogate spiking solution. Transfer 2mls of hexane to vial, cap and shake vigorously for 5 minutes. The hexane layer (top) can now be transferred to a vial for cleanup or analyzed directly. Results must be divided by 15 due to the concentration factor of the extraction procedure.

If sample requires cleanup procedure, transfer 1-3ml of extract to a Supelco cleanup cartridge that has been pre-rinsed with 2-ml hexane. Discard the first 0.5ml of extract that elutes from the cartridge and then collect the remaining fraction in a small vial. The extract is now ready for analysis by direct injection into the gas chromatograph.

Section 8
Revision 1
Date 1/1/93
Page 12 of 16

<u>OA/OC Analyses</u>. For each set of sample analyses, a method blank is to be analyzed. A method blank consists of the anhydrous sodium sulfate shaken with 10ml of hexane and spiked with 10 μ l of surrogate spiking solution in a VOA vial. If sample cleanup was performed on any of the samples analyzed, a method blank must also be analyzed in which the method blank was passed through a Supelco Florisil cartridge prior to analysis.

For pesticide analysis a 25 ppb standard is to be analyzed to confirm the calibration curve at the beginning of each 12 hour shift. The measured concentration for each pesticide must be within 80% - 120% of the true value for the calibration to be considered still valid. If the calibration is determined not to be valid, a 4-point calibration is to be performed using the following pesticide concentrations, 5.0, 10.0, 25.0 and 50.0 ppb.

After each 10 samples are analyzed, a pesticide/PCB matrix spike and matrix spike duplicate are to be extracted and analyzed. If any samples were found with PCB's present, the PCB used in spiking should be the type that is in the samples. For soils the recommended spiking concentration is 1 ppm for PCB's and 20 ppb for pesticides. This is performed by spiking 10 +/- .1 gram of sample with 50ul of 200 ppm PCB stock in methanol, or with $10\mu l$ of the 20 ppm pesticide stock solution. The spiked sample is then mixed well with 10ml of hexane for 2 minutes and then sonicated for 10 minutes. For waters the recommended spiking concentration is .1 ppm for PCB's and 2 ppb for pesticides. This is performed by spiking 30 +/- .1 gram of sample with 15 ul of 200 ppm PCB stock or with 10 μl of the 2 ppm pesticide stock solution. The spiked sample is then mixed well with 2ml of hexane for 5 minutes. After analysis of the matrix spike and matrix spike duplicate, the percent recovery is to be determined as follows:

The acceptable % recovery limits are 65% to 125%. In addition to determining the percent recovery, the % RPD is to be determined and reported. To calculate the % RPD use the following calculation:

$$\% RPD = \% Rec_{MS} - \% Rec_{MSD}$$

$$\frac{}{(\% Rec_{MS} + \% Rec_{MSD}/2)}$$

This value is to be reported along with the % Recovery of the matrix spike and matrix spike duplicate.

Section 8 Revision 1 Date 1/1/93 Page 13 of 16

187278

<u>Data Evaluation and Pesticide/PCB Quantitation</u>. For pesticide analysis, dual columns are to be used, one for quantitation (Rtx-5) and the other (DB-17) for confirmation. Upon reviewing the data obtained from the quantitation column, any "hits" obtained on this column <u>must</u> be confirmed on the other column before the result is accepted as a true hit. The quantitation values must also be within 1 to 10 times of each other for the confirmation to be considered valid.

The first step in PCB analysis is to identify the type(s) of PCB present in the sample analyzed, if any. Each of the PCB's have a characteristic chromatogram fingerprint that can be used to identify them. Some of the PCB's have peaks appearing at the same retention time but the integrated area of these peaks relative to other peaks is different. The analyst must become experienced in using these relative peak ratios in the identification of the PCB. The use of transparent overlays prepared of each PCB type is used by TEG chemists and is very useful for identification purposes. Caution must still be used, however, when analyzing samples that have more than one type of PCB present.

Once the PCB type has been identified the 1 ppm standard of that specific PCB is to be analyzed to confirm the identification of the PCB. After the standard has been run, quantitation can then be performed. If the sample if free from other contaminants, the total area of the PCL may be used. A ratio of the total peak area of the PCB in the sample can then be made to that of the 1 ppm PCB standard to calculate the total PCB present in the sample. For example, if a sample has PCB 1254 present and the analysis resulted in a total area count of 5680, the concentration of this PCB can be calculated as follows:

Total area PCB 1254 1 ppm standard = 4850.63

$$\frac{1 \text{ ppm}}{4850.63} = \frac{X}{5680}$$

This ratio calculates "X" as the concentration of PCB 1254 in the sample to be 1.17 ppm.

If there are other contaminants present in the soil sample, which happens in most cases, the concentration of PCB present is to be calculated using the total area of a minimum of 3 peaks that have been used to identify the PCB. The total area of these peaks is then compared to the total area of these same peaks of the 1 ppm PCB standard and the concentration of the PCB calculated as indicated above.

The recommended peaks to use for each of the PCB's and their retention times are listed below. Most of the PCB's have only three peaks listed. For those PCB's which are more difficult to identify, there are more times listed.

PCB TYPE	PEAK RETENTION TIMES 18727
1016	10.4, 11.1, 12.4
1221	5.02, 7.90, 8.58
1232	8.55, 10.33, 12.26
1242	10.4, 11.1, 12.4, 20.86, 22.11
1248	12.3, 16.7, 17.55
1254	20.88, 21.98, 23.25
1260	21.85, 23.10, 26.2

8.8 EPA METHOD 604/8040

Introduction. This method can be used in determining the concentration of Phenol's present in soil and water samples. This procedure involves extracting the Phenol's from a 10 gram sample of soil or a 30 gram sample of water using freon. The extract is then analyzed by gas chromatography using an flame ionization detector. The detection limits for each Phenol using this procedure are listed below.

Calibration. Calibration may be accomplished by either internal or external standard techniques. Standards are prepared at three or five concentrations spanning the linear range of the instrument. The standards are then analyzed and the response factor evaluated for linearity. If linearity is not achieved a second set of analyses is run. If linearity is still not achieved, a second set of standards is prepared and the process repeated. Linearity is acceptable when response factor values are within 15% standard deviation for internal standard calculations or if the mean residual from the linear regression is within 15%.

Apparatus and Reagents.

- Freon gas chromatographic grade
- Sodium sulfate anhydrous
- 5 N Sulfuric acid solution
- Phenol stock solutions 2000 ppm in 2-propanol or other suitable solvent.
- Surrogate spiking solution 1000 ppm of 2-Fluorophenol in 2-propanol.
- Silica gel cartridges prepared with activated 100/200 mesh silica gel.
- VOA vials 40mL glass w/polyethylene screw caps.
- Shimadzu 14A GC with an flame ionization detector using a PTE-5 QTM, 15 meter, 0.53 mm diameter & 0.5 μm film column and Helium as carrier gas.

Injection port temp - 175 degrees C Detector temp - 225 degrees C Column temp - 65 degrees C/3 min. to 185 degrees C at 10/min 3 μl injection for analysis

Section 8
Revision 1
Date 1/1/93
Page 15 of 16

Sample Preparation. Tare a 40mL VOA vial on a top loading balance and transfer 10 gram +/-.1 gram of soil sample in the vial. Add enough anhydrous sodium sulfate to vial until the soil becomes free flowing. Spike the soil sample by adding $10 \mu l$ of the surrogate spiking solution directly into the soil. Add 10ml of freon and cap vial. Shake sample well for 2 minutes and then sonicate for 10 minutes.

Sample can now analyzed by direct injection into the gas chromatograph.

For water samples, place 30.0 grams of water to be analyzed in a voa vial. Add enough acid to water to bring pH to below 2. Spike the water sample by adding 3 μ l of the surrogate spiking solution directly into the water. Transfer 2mls of freon to vial, cap and shake vigorously for 5 minutes. The freon layer (bottom) can now be analyzed directly. Results must be divided by 15 due to the concentration factor of the extraction procedure.

If any extract requires cleanup procedure, transfer 1-2ml of the extract through a silica gel column that has been prewashed with 2-3ml of freon. Discard the first 0.5ml of the extract that elutes from the cartridge, then collect the remaining fraction in a VOA vial. The extract is now ready for analysis by direct injection into "e gas chromatograph.

QA/QC Analyses. At the beginning of every 12 hour shift a continuing calibration is be run to verify that the calibration is still valid. The recommended concentration of this standard is 1 ppm of each component. The determined concentration of each phenol must be within 80% to 120% for the calibration to be considered valid. If these limits are not met a 4 point calibration is to be performed to reestablish calibration. The standard concentrations that are recommended are 0.1, 0.50, 1.00, and 10.0 ppm for both water and soil analysis.

For each set of sample analyses, a method blank is to be analyzed. A method blank consists of the anhydrous sodium sulfate shaken with 10ml of freon in a VOA vial. If sample cleanup was performed on any of the samples analyzed, a method blank must be analyzed in which the method blank was passed through a prepared silica gel column.

After each 10 samples are analyzed, a Phenol matrix spike and matrix spike duplicate are to be extracted and analyzed. The recommended concentration to spike soil samples is 1 ppm. This is performed by spiking 10 + /- .1 gram of a soil sample with $15 \mu l$ of the 2000 ppm phenol stock solution. The recommended concentration to spike water samples is .2 ppm. This is performed by spiking 30 + /- .1 gram of a water sample with $3 \mu l$ of the 2000 ppm phenol stock solution. The spiked sample is then extracted according to the procedure outlined above. After analysis of the matrix spike and matrix spike duplicate, the percent recovery is to be determined as follows:

Section 8
Revision 1
Date 1/1/93
Page 16 of 16

% Recovery = Conc. determined X 100 Conc. Spiked

187281

The acceptable % recovery limits are 65% to 125%. In addition to determining the percent recovery, the % RPD is to be determined and reported. To calculate the % RPD use the following calculation:

$$\% RPD = \% Rec_{MS} - \% Rec_{MSD}$$

$$(\% Rec_{MS} + \% Rec_{MSD}/2)$$

This value is to be reported along with the % Recovery of the matrix spike and matrix spike duplicate.

8.9 REAGENT STORAGE

Reagents used in analysis are stored as described in Table 8-1.

TABLE 8-1. REAGENT AND STANDARD STORAGE

Chemical

Method of Storage

Methanol: Freon-113:

In cabinet separate from other solvents In cabinet separate from other solvents

Neat Standards:

Refrigerator away from samples

Working Standards:

Methanol standards in refrigerator away from samples

Working Standards:

F113 standards in separate refrigerator

Reagent bottles are labeled with the date received, date opened, and recorded on a solvent and standard log sheet.

Section 9 Revision 1 Date 1/1/93 Page 1 of 10

9.0 CALIBRATION PROCEDURES AND FREQUENCY

187282

9.1 INSTRUMENTATION LIST

A list of TEG's primary instrumentation and applicable methodology is summarized below in Table 9-1.

TABLE 9-1 INSTRUMENTATION LIST

INSTRUMENT	DETECTORS	<u>METHODS</u>
Gas Chromatograph Buck HC-404	FID, PID, Hall, ECD IR	8010, 8020, 8015, 8100, 8080 418.1
Tekmar LSC 2000	Purge and Trap	5030

9 > STANDARD RECEIPT AND TRACEABILITY

Sundards and solvents used in the preparation of substocks and analyses are recorded on receipt in the TEG Standard and Solvent Log-in Form, Figure 9-1. Bottles are labeled with the date received and opened. Each standard and solvent is "blank" tested for impurities prior to usage and periodically before each substock or working standard is prepared. Each type of solvent used by TEG is stored in a cabinet separate from the others. Each analytical group (and matrix) of standards is stored in refrigerators separate from other standards. Neat standards are similarly kept in separate, respective refrigerator compartments. Additional storage information is presented in Section 8.9.

9.3 STANDARD SOURCES AND PREPARATION

Preparation of standards and substocks is recorded on the Standard Preparation Log, Figure 9-2, according to procedures listed in the analytical methods, (Section 8).

Section 9 Revision 1 Date 1/1/93 Page 2 of 10

Figure 9-1. Solvent and Standard Log-in Form

TEG SOLVENT AND STANDARD LOG-IN FORM

TE	INITIALS	P.O. NUMBER	SCLVENT	GRADE	VENDOR	BATCH NUMBER	BLANK TEST	COMMENTS
-								
							•	
						i		
					_			
				-			<u>.</u>	
	<u> </u>					! 	<u> </u>	<u> </u>
					_			
	<u> </u>				_			
_								

Section 9 Revision 1 Date 1/1/93 Page 3 of 10

Figure 9-2. Standard Preparation Log

TES STANDARD PREPARATION LCG

PREPARATION DATE	INITIALS	DESCRIPTI	CN	ANALYTE	STARTING STANDARD OR SOLVENT	HIZZIA	DILUTION FACTOR	FINAL CONG.
	!							
	i					1	<u> </u>	
	-					-	<u> </u>	1
	<u> </u>	!		<u> </u>		<u> </u>		!
		j					1	!
	1		!			:		
	!		<u>:</u>				:	!
	!			1		-	<u>: </u>	!
			į	:			:	
				<u> </u>		!		1
	<u> </u>		_ <u>-</u> -	<u>:</u>			<u>. </u>	
						:		
	!			1		:		
	!			<u> </u>				

Section 9 Revision 1 Date 1/1/93 Page 4 of 10

9.4

INSTRUMENTATION CALIBRATION

9.4.1 Instrument Group Calibration

Table 9-2 has been prepared to summarize specific routine calibration procedures for each instrument group.

9.4.2 Initial Calibration and Linearity

Calibration standards are prepared for three to five concentration levels depending upon the method for use in determining the linearity of the system. The linearity of the response to concentration is assessed over the working range of the standards which are prepared for each analyte. A low level standard is prepared at or near the method detection limit. The medium and high level standards are prepared to correspond with the range of concentrations anticipated in the working samples and within the range of the detector. A solvent blank is run prior to the initial calibration runs in order to provide information regarding system cleanliness.

If the relative standard deviation of the linearity data is less than fifteen percent (15%), linearity is assumed and the average response factor, as defined by the slope of the response curve, is used in calculating concentration of unknown samples. If the relative standard deviation is greater than fifteen percent (15%), a calibration curve is prepared for each compound and the response factor for each sample is determined directly from the calibration curve. All calibration procedures and results are entered into the logbook for the instrument (see Instrument Initial Calibration Log, Figure 9-3, and Continuing Calibration Form, Figure 9-4).

Section 9 Revision 1 Date 1/1/93 Page 5 of 10

TABLE 9-2 INSTRUMENT CALIBRATION

Free	Daily + 10 Samp.	Daily + 10 Samp
Accept/Reject Criteria-Cont.	< 15%RPD	< 15%RPD
# Standards Cont. Cal.	I + Dup.	l + Dup.
Frequency	Cont. Cal Failure	Cont. Cal Failure
Accept/Reject <u>Criteria-lait.</u>	Linear Reg. < 15% RPD Response Factor	Linear Reg. <15% RPD Response Factor
// Standards Init. Cal.	m	€
Standard Source	Supeko Chemserve	In-House Chemserve
Instrument	Gas Chromatograph	x

Section 9 187287 Revision 1 Date 1/1/93 Page 6 of 10

Figure 9-3. Instrument Initial Calibration Log

CALIBRATION LOG

DATE: ANALYST: INSTRUMENT:

ANALYTE	FINAL CONC.	ABSORB.	PEAK AREA	CONC. OF STK. USED	VOL.OF STK. USED	FINAL VOLUME	SOLVENT
						<u> </u>	
					<u> </u>		
				+			
				+ -			
	<u> </u>			<u> </u>		•	
			 				
				1			

CORRELATION:

X COEFFICIENT:

STD. ERROR:

Section 9
Revision 1
Date 1/1/93
Page 7 of 10

Figure 9-4. Continuing Calibration Form

						DATE:
Septu	= 0 Co1:	umn [Inj.	, Col, Det	Temp [Gas Fle	ow & Pressure Sig	nature
TIME	PROJECT NUMBER	ANALYST	STANDARD NUL TR	CONCENTRATION	RESPONSE FACTOR	RELATIVE PERCENT DIFFERENCE
		1		·		
				<u> </u>		
		ļ_				
				1		

9.4.3

QC Check Standards

On completion of the initial calibration, a QC check standard is run. The QC check standard is an independent check standard to validate the initial calibration.

9.4.4 Continuing Calibration

A continuing calibration check is run using either a standard prepared similarly to the curve standards or an EPA or certified commercial NBS-traceable check standard to validate the calibration. Following a solvent blank, a continuing calibration standard is run before starting each day and every ten samples, or every batch, if less than ten samples, to validate response factors and retention times. The percent difference of the check response factor to the initial calibration response should be less than 15%. Retention time calibration checks should fall within 5% of the Initial calibration. The results of the calibration checks are also entered on the Calibration Log for the instrument. If the check standards fail the continuing calibration criteria, initial calibration procedures must be implemented and continuing calibration rerun.

Surrogate standards and matrix spikes are used in analyses in order to monitor recovery of each sample through the extraction procedure. A separate log of extractions and recovery is mantained for each method. Absolute recovery varies with the method and the matrix. Recovery criteria are presented in Table 5-1.

9.4.5 Equipment Monitoring

Balances. Laboratory balances are to be calibrated prior to each group of weighings, or daily, according to the following protocol:

- check level of balance;
- zero the balance:
- weigh standard weight; and
- record true weight, measured weight, and error in Balance Calibration Log and sign.

<u>Refrigerators</u>. The purpose of monitoring refrigerator temperatures is to maintain a record of temperature variation which may affect sample integrity. Each working day, the refrigerator temperature will be monitored from the thermometer mounted in the unit and entered on the Refrigerator Temperature Log (Figure 9-6). Should the temperature be unacceptable, the QA Manager will be notified immediately.

Mobile Laboratory Temperature. The purpose of monitoring refrigerator temperatures is to maintain a record of temperature variation which may affect sample integrity. Each working day, the refrigerator temperature will be monitored from the thermometer mounted in the unit and entered on the Refrigerator Temperature Log (Figure 9-6). Should the temperature be unacceptable, the QA Manager will be notified immediately.

Section 9 Revision 1 Date 1/1/93 Page 9 of 10

Figure 9-5. Balance Calibration Log

TEG Mobile Lab

Balance Calibration Log

Cate	Initial	Reference Wt. 1	Reference Wt. 2
	1		
_			
		1	
	1	j	1
	1		
	1	i	
			1
_		1	!
	- i		1
	1	Ti Ti	
	1	1	!
	1	1	
	1		1
		i	
		1	
	1	1	
	1		
	1		
	-i		
-		1	
		1	
-	<u> </u>		
		1	1
	7		
	1		

187293 Section 9 Revision 1 Date 1/1/93 Page 10 of 10

Figure 9-6. Refrigerator Temperature Log

		REI	RIGER	TE	З С,	IE: I:	IPE	:R:	\=:	R.E	: :	.cg								
Re	Refrigerator Period to								_											
DATE	TIME	ANALYST	INIT	RE	FRI De	ig:	RA ee	TC S	З Се	:: :::	MP ig	IP.	ÀT de	UR.	E		AC	TIC:	1	
				3	1	2	3	÷	5	કં	7	3	9	10						
					1	Ţ	Ī	I		I	Ī									
					• •	Ī	1	İ	1	:	Ī		İ							
					!	į		i			\prod						_			
					į	<u> </u>		-	İ							_				
	<u>-</u>		<u>l i</u>	_	<u>i</u> .		<u> </u>	<u> </u>	<u>i</u>		!	<u> </u>								
					:	<u> </u>	1	_		<u> </u>		<u> </u>	<u> </u>							
	-			1		<u>i</u>	<u> </u>	_	<u> </u>	-	_	<u> </u>								
			1 1			Ļ	<u> </u>	<u> </u>												
			<u> </u>		:	1	<u> </u>	_	<u>i </u>	_										
	<u> </u>		1 1		<u>:</u>	<u> </u>	<u> </u>	<u> </u> _	!	<u> </u>										
				1	1	1	1	•	1				_	_						
			+	-	-	<u> </u>	<u> </u>	!	1				_	_ !						
<u> </u>			<u> </u>	1	i i	<u>!</u>	1	<u> </u>	:				_							_
-	 		<u> </u>	<u> </u>	<u>:</u>	<u> </u>	<u>i </u>	_			j	-	1	<u> </u>	_					
<u> </u>		<u> </u>			<u>i</u>	<u> </u>	! !	<u>-</u>		1	-!	- [-	4				_		
				1		1	<u> </u>			!	<u>i</u>	<u> </u>	<u> </u>	<u> </u>		_				
			<u> </u>	<u> </u>	<u>i</u>	\vdash			1	-	<u> </u>	<u> </u>	1	1		_				
				1	-	╁	<u> </u>	_ ;	<u> </u>	1	<u> </u> 	1	+	<u>1</u>						
- 					1		<u>.)</u> 	<u>;</u>	i	<u> </u>	+	\dashv	+	$\frac{1}{1}$						
		- <u></u> -		Ť	Ė				+	\dashv	-	1	\dagger	\dagger						
	<u>-</u> :	- '	- i	i			i	-	T	\dashv	+	T	7	┪			_			
	<u> </u>		i	İ				1	i	T	Ť	\forall	+	\dagger				_		_

Section 10 Revision 1 Date 1/1/93 Page 1 of 3

10.0

PREVENTIVE MAINTENANCE PROCEDURES AND SCHEDULES

10.1 ROUTINE MAINTENANCE

The preventive maintenance program combines both in-house procedures and maintenance provided by the manufacturers and vendors of the equipment. Major equipment and components are periodically inspected and tested by manufacturers representatives, as appropriate. Routine parts replacement and system modifications are performed by qualified personnel. Listed in Table 10-1 are routine maintenance activities for the equipment listed in Section 9.

TABLE 10-1. MAINTENANCE ACTIVITIES

<u>Instrument</u>		Activity		Frequency
Gas Chromatograph	Check	Septum Change Septum Check Carrier Clean PID lamp Leak Check	Daily	As required Daily As required Daily
IR Spectrophotometer		Clean Cell & Extractor		Each use
Balance		Clean, Calibrate		Daily
Refrigerator		Clean, Calibrate		Daily
Generators		Oil Change, Filters		100 Hours
Vehicle Engine		Oil Change, Filters		3000 miles

10.2 DOCUMENTATION

A maintenance logbook is kept to record problems, routine maintenance, and corrective service for each instrument (Figure 10-1). Items which require regular maintenance, such as filters, compressors, generators, and engines, are labeled with the date of last maintenance and due-dates (or criteria) for next regular maintenance.

Section 10 Revision 1 Date 1/1/93 Page 2 of 3

10.3 CONTINGENCY PLAN

A stock of key replacement items is maintained so that down-time is reduced. Key personnel are trained in ares of routine maintenance, trouble shooting, and repair of electronic and mechanical failure. This is especially critical in the TEG mobile laboratory operation as backup equipment and factory maintenance may not be readily available.

In the event of total equipment failure, or the inability to repair equipment on-site, TEG will arrange either for back-up equipment and parts to be delivered or for service personnel to effect repairs as needed. If a substantial down-time is anticipated, judgement will be made whether to store samples until repairs are completed or to transfer samples to another TEG facility for analysis.

Section 10 Revision 1 Date 1/1/93 Page 3 of 3

Figure 10-1. Repair Maintenance Log

:					

TEG REPAIR MAINTENANCE LOG

DATE	INITIALS	DESCRIPTION OF PROBLEM	DESCRIPTION OF REPAIR	DATE REPAIRED/WHCM
		<u> </u>		
	1			
	1			
	1			

Section 11 Revision 1 Date 1/1/93 Page 1 of 3

11.0 QC CHECKS, PRECISION AND ACCURACY AND DETECTION LIMITS

11.1 LABORATORY QC CHECKS

11.1.1 Equipment QC Checks

Listed below in Table 11-1 are equipment monitoring activities with respect to QC checks. Monitoring of this equipment is documented on forms presented in Section 9.

TABLE 11-1. EQUIPMENT MONITORING ACTIVITIES

<u>Instrument</u>	<u>Activity</u>	<u>Frequency</u>
Gas Chromatograph	Ongoing Calibration	Daily
IR Spectrophotometer	Calibration	Each use
Balance	Caribrate	Daily
Refrigerator	Monitor Temperature	Daily
Mobile Laboratory	Monitor Temperature	es Daily

11.1.2 Chemistry QC Checks

Several types of QC checks are standard to TEG analyses. The standard operating procedures for each analysis have specific types and frequency of QC checks which are followed at a minimum. In general, TEG QC checks include an evaluation of:

- blanks;
- precision;
- accuracy;
- recovery; and
- method detection limits.

If a method does not specify minimum QC checks or if the method QC requirements are less stringent than those listed in below in Table 11-2, the laboratory will use the QC guidelines presented in Table 11-2.

Section 11 Revision 1 Date 1/1/93 Page 2 of 3

TABLE 11-2 CHEMISTRY QC CHECKS

BLANKS

Method Reagent Blanks - prepared and analyzed at greater of one per sample set or 5%

MATRIX SPIKES - prepared and analyzed at greater of one per sample set or 5%

- minimum of one set per matrix

QC CHECK SAMPLES - analyzed blind in duplicate semi-annually

QC CHECK
STANDARDS

- analyzed at greater of one per sample set or 5%

DUPLICATE SAMPLES - Duplicate samples or duplicate matrix spikes will be analyzed

at greater of one per sample set or 5%; minimum of one per

matrix

CONTINUING CALIBRATION

STANDARDS - Same as, or in lieu of QC check standards

11.1.3 Additional QC Checks

A calculated recovery of sample during analysis is accomplished by the addition and subsequent analysis of surrogate or matrix spiking compounds. Surrogate spiking compounds, which serve as substitute analytes, differ for each analysis and are usually not naturally occurring compounds. The surrogates are added to samples for analysis after aliquots of the sample have gone through the preparation process. Surrogates are chosen such that they are chemically similar to the analytes, with similar response, but do not interfere with determination of the analytes.

11.2 PRECISION AND ACCURACY

Precision is determined through evaluation of percent difference in duplicate analysis of samples, standards and surrogates. Precision may also be measured by evaluating the standard deviation of multi-point calibrations or the standard deviation of the high point in a span calibration.

Section 11 Revision 1 Date 1/1/93 Page 3 of 3

Precision, as determined through percent difference in duplicate analysis of samples, standards and surrogates, is calculated as:

Precision =
$$\frac{A - B}{A + B} \times 100$$

where "A" is the larger value and "B" is the smaller value of two replicate analyses.

Accuracy, as percent recovery, is calculated from analysis of surrogate and matrix spike samples as follows:

Accuracy =
$$\frac{A - B}{C} \times 100$$

where "A" is the analyte determined experimentally from the spike sample; "B" is the background level by separate analysis of the unspiked sample; and "C" is the amount of spike added.

11.3 METHOD DETECTION LIMITS AND PRACTICAL DETECTION LIMITS Method detection limits (MDLs) are derived from in-house data according to the guidelines of 40CFR 136. Replicate samples or standards are analyzed, with the method detection limit being based on 3 times the standard deviation for the analyses.

Practical quantitation limits (PQLs) are defined as ten times the standard deviation resulting from the calculation used in determination of method detection limits.

MDLs and PQLs are re-evaluated every 12 months by in-house analyses or each time a modification is made to an analytical procedure.

Section 12 Revision 1 Date 1/1/93 Page 1 of 1

12.0 DATA REDUCTION, VALIDATION AND REPORTING

12.1 DATA REDUCTION

Each analysis is recorded by the field chemist on an Analysis Log (Figure 7.3) for the instrument being used. TEG uses microcomputers for data storage and reporting purposes. Initial chromatographic data gathering is performed either by integrators or computers which print a chromatogram and report integration. Each chromatogram is labeled during analysis to identify the project and sample number. It is the responsibility of the field chemist to evaluate each chromatogram with regard to correct integration, peak identification, and calibration. Verified numeric data from the chromatogram is then manually entered by the analyst into a spreadsheet on the microcomputer. A spreadsheet file, with a unique file name related to the project number and analysis, is created from a generic starter file for each project and analytical method. At the end of each day, the active file is transferred to a backup floppy. This backup floppy is stored in the final job file.

12.2 DATA VALIDATION

At the end of each day the field chemist (analyst) inspects the extraction logs, analysis logs, calibrations, raw data, and calculated data. The printed spreadsheet is placed in the project file and submitted to the QA officer for review. The quality assurance officer then reviews the data for errors and makes a determination as to whether each quality control parameter is within the limits set for the analysis. If the data are acceptable, they are approved for reporting. Both the analyst and QA officer are responsible for review of these measures. In the event that a QC parameter is unacceptable, corrective action will be taken. Corrective actions are discussed in Section 13.

12.3 DATA REPORTING

TEG uses computer automation where feasible to reduce potential for error between data reduction and reporting. Data reporting is performed following review of data and calculations by the analyst. The data generator prepares a preliminary report for review by the QA Manager. The QA Manager then reviews the report and prepares a QA summary. The generated report is then reviewed for typographical errors and forwarded to the laboratory director for review, signature and release to the client. A sample report is enclosed in an appendix.

12.4 DATA STORAGE

Hard copies of the final printout and report are maintained in the project file by job number along with printed chromatograms, logsheets, chain of custody records, floppy disk, and data tables. The purpose of this document control procedure is to provide guidelines to reduce the potential for document loss and mix-ups. Both electronic and printed media records are stored for a minimum of two years.

Section 13 Revision 1 Date 1/1/93 Page 1 of 3

13.0 CORRECTIVE ACTION

13.1 CRITERIA

Whenever the quality control goals set for precision or accuracy of data are not achieved, as summarized in Table 13.1, a program of corrective action is initiated.

13.2 RESPONSIBLE INDIVIDUALS

The first step in corrective action is identification of the source of the problem. Initial identification responsibility is with the analyst who will spot most problems. The quality assurance officer is responsible for identification of problems which the analyst may have overlooked. He must also initiate the corrective action and review the effectiveness of the action. The laboratory director is responsible for review of the QA Manager's reports.

13.3 CORRECTIVE ACTIONS

When a quality control problem is noted, the following steps are taken to identify and correct the problem:

- the hard copies of the data are re-examined;
- the analyst re-analyzes the sample(s), as appropriate;
- if the problem is not resolved by analysis, the QA Manager or the laboratory director is consulted to provide additional information about rectifying the problem; and
- if the problem cannot be solved in-house, equipment repair contractors, manufacturer's representatives, or outside consultants are contacted, as necessary to correct the problem.

Section 13 Revision 1 Date 1/1/93 Page 2 of 3

TABLE 13-1 SUMMARY OF CORRECTIVE ACTION PROCEDURES (applicable to GC and IR analyses)

OC ACTIVITY	ACCEPTANCE CRITERIA	RECOMMENDED ACTION
System Blank	Response < MDL	Repeat, if same, identify source eliminate source
Reagent Water Blank	Response < MDL	Repeat, if same, identify source, correct problem
Reagent Blank	Response < MDL	Repeat, if same, identify source, Replace reagent
New Standard Validation	Relative % difference < 15%	Re-analyze; if still unacceptable, Reject Standar ⁴
Method Blank	Response < MDL	Repeat, if still unacceptable, identify cause, re-process, Re-analyze previous sample set
Initial Calibration	Relative % difference < 15%	Re-analyze; if still unacceptable, make fresh standards
QC Check Standards	Relative % Difference < 15%	Re-analyze; if still unacceptable, make fresh standard, use new standard
Cont. Cal. Standards	Relative % Difference < 15%	Re-analyze; if still unacceptable, make fresh standard, use new standard, Re-analyze previous sample set
Matrix Spikes	Method Control Limits	Re-analyze, if extremely low recoveries.

13.4 EXTERNAL SOURCES

Corrective actions are also initiated when other problems or irregularities are observed during external auditing procedures, client review, or by regulatory review. These externally initiated corrective actions may be a result of performance evaluations, audits, split samples or standards submitted by clients, or other certification procedures.

Section 13 Revision 1 Date 1/1/93 Page 3 of 3

13.5 NOTIFICATION OF PERSONNEL

The field chemist and QA Manager will initially discuss the identification of problems and implementation of corrective actions. If the problem is of a routine nature the QA Manager will make note of it in his QA report for the project to the laboratory director. If the problem is of a severe nature, the situation will be discussed immediately with the laboratory director so that he may be involved in the decision-making process regarding corrective action.

Section 14 Revision 1 Date 1/1/93 Page 1 of 3

14.0 PERFORMANCE AND SYSTEM AUDITS

14.1 GENERAL REQUIREMENTS

Performance and system audits are achieved through both external and internal processes. These audits are conducted with specified frequency to review and evaluate the individual components and the overall measurement system with regard to proper selection and usage. The following sections include descriptions and frequency of both internal and external audits.

14.2 SYSTEM AUDITS

14.2.1 System Audit Scope

The system audit is designed to evaluate the individual components of the operational system. A determination is made regarding the proper function, proper methodology, and relationship to the CompQAP. Resulting output from the system audit indicates deficiencies and areas which need improvement. Listed on the following page (Figure 14-1) are the system components which are evaluated through the system audit.

14.2.2 Internal System Audits

Internal system audits are conducted annually at each TEG facility. The audits are conducted by either the TEG corporate QA Manager or a laboratory director from another TEG facility. Each system component listed on the Internal Audit System Audit Form (Figure 14-1) is assessed and graded during the procedure.

14.2.3 External System Audits

TEG welcomes external system audits from regulatory agencies clients and peer groups. The external system audit provides an opportunity to improve the TEG system operation in ways that may be different from the internal audit and to evaluate the laboratory with respect to other laboratory operations.

14.3 PERFORMANCE AUDITS

14.3.1 Performance Audit Scope

Performance Audits are conducted for the purpose of evaluating the analytical capability of the laboratory. Both the accuracy of the overall analysis and the individual portions are evaluated. The performance audit consists of each of the following: blind samples, split samples, quality control samples, regulatory or commercial check samples, and blind spikes.

Section 14 Revision 1 Date 1/1/93 Page 2 of 3

14.3.1 Internal Performance Audits

Internal performance audits are conducted semi-annually by the corporate QA Manager. Blind samples, QC samples and blind spikes are prepared at a different TEG facility. Sample splits are forwarded to another TEG laboratory for analysis. Commercial check samples may also be obtained. The report of analysis is prepared in the same manner as normal analytical report. The QA Manager reviews this report in comparison with known values and results of split analyses. The QA Manager then makes a formal report of the audit to the laboratory director.

14.3.2 External Performance Audits

External performance audits are requested by some regulatory agencies and clients. TEG conducts performance audits as required by regulatory agencies and clients.

Section 14
Revision 1
Date 1/1/93
Page 3 of 3
187304

Figure 14-1. Internal System Audit Form

rigule 14-1. Internal system	וו הע	dit Porm
 SAMPLING PROCEDURES		
 Sample Containers		
 Cleaning		
 Waste Disposal		
ALVELE GUSTORY		
 SAMPLE CUSTODY		
 Field Custody		
 Laboratory Custody		
ANALYTICAL PROCEDURES		
 Method References		
Waste Disposal		
•		
 CALBRATION		
 Instrumentation List		
 Standard Receipt and Traceability		
 Standard Sources and Preparation		
 Instrument Calibration		
DREVENTATIVE MAINTENANCE		
 PREVENTATIVE MAINTENANCE Routine Maintenance		
 Documentation		
 Contingency Plans		
 Contingency Fians		
QUALITY CONTROL CHECKS		
• • • • • • • • • • • • • • • • • • • •		
Equipment QC checks		
Chemistry QC checks		
 Additional QC checks		
 Precision and Accuracy		
 Method and Practical Detection Limits		
DATA DEDUCTION WALKDAMON AND DEDOCTOR		
 DATA REDUCTION, VALIDATION AND REPORTING Data Reduction		
 Data Validation		
 Data Reporting		
 Data Storage, Document Control		
 Data Glorage, Document Control		
CORRECTIVE ACTION		
 Criteria		
 Responsible Individuals		
Corrective Actions		
External Sources		
 Notification of Personnel		EXPLANATION
DEDEGRAVA AND AVAILABLE AND AVAILABLE		.
 PERFORMANCE AND SYSTEM AUDITS	_	Passing
 General Provisions	2.	
 System Audits Performance Audits	3.	Unacceptable
 remorinance Audits		
OUALITY ASSURANCE REPORTS		

15.0 QUALITY ASSURANCE REPORTS

187305

QA reports are prepared by the QA Manager on an annual basis. These reports contain:

- the results of performance audits;
- summary of individual project QA;
- periodic assessment of accuracy, precision and detection limits;
- significant QA/QC problems and recommended solutions; and
- the results of corrective actions implemented.

The QA Manager's report is prepared and delivered to the laboratory director and president of TEG. Overall laboratory performance and the results of system and performance audits are reviewed by the laboratory director and president of TEG. External QA reports are prepared, as required, by regulatory agencies and clients.

16.0 RESUMES 187306

SHERI WILDER HARTMAN, M.A.

President, Transglobal Environmental Geochemistry, Inc. Co-Founder, Solana Beach, California

Education:

Masters Program, Geology/Geochemistry - 1978-1980; Tulsa University,

Tulsa, Oklahoma

M.A., Creative Arts - 1976; Purdue University,

West LaFayette, Indiana

B.A., Chemistry and Biology - 1976; Purdue University,

West LaFayette, Indiana

Summation of Employment History:

1990 to Date: Transglobal Environmental Geochemistry, Inc., Solana Beach, California,

Geochemist & QA Manager

1988 to Date: Transglobal Exploration & Geoscience, Inc., Solana Beach, California,

President and Co-Founder

1980 to 1987: Unocal Science and Technology Division, Brea, California,

Research Geochemist

1976 to 1980: Dept of Energy, Energy Technology Center, Bartlesville, Oklahoma

Research Chemist

Professional Experience:

Ms. Hartman's professional career includes over thirteen years of varied experience in petroleum and environmental geochemical applications. In addition to extensive laboratory analytical work, she has been engaged in a variety of field data collection programs. Ms. Hartman has served as director of numerous oceanographic and land-based geochemical programs with Unocal Research and Transglobal Exploration. With TEG, Ms. hartman is responsible for field sample collection and analysis, reporting, and management of the Quality Assurance Program.

President and Co-Founder:

Transglobal Exploration & Geoscience March 1988 to Present

Primary Duties:

As President and Co-founder of Transglobal & Geoscience in 1988, Ms. Hartman conducts all the duties of a president and is a senior geochemist experienced with gas chromatographic systems and data interpretation.

Research Geochemist:

Unocal Science and Technology December 1980 to June 1987

187307

Primary Duties:

Direct all geochemical programs for Unocal Indonesia, Latin America, Dutch North Sea, and Egypt; provide technical interpretation and reports.

Participate in and provide technical support for marine geochemical exploration programs for petroleum in the Southern California Borderlands (1981), offshore California (1981), Behring Sea (1983) and North Sea (1985).

Direct surface-geochemical surveys for petroleum exploration in Nevada, Oregon, and Alberta, Canada.

Organize and conduct Basin Modeling Short Course for Unocal personnel.

Development of new HPLC methods for characterizing chemical-compound classes in petroleum.

Initial organization of geochemical database.

Research Chemist:

Department of Energy, Bartlesville Energy Technology Center September 1976 to December 1980

Primary Duties:

Organize geochemical studies to enhance on-going oil-to-oil correlation studies.

Development of new correlation methods using isotope ratio data.

Conduct petroleum studies using LC, HPLC, IR, AA and MS data.

Direct petroleum analysis lab.

Section 16 Revision 1 Date 1/1/93 Page 3 of 11

BLAYNE HARTMAN, Ph.D.

187308

Vice President, Transglobal Environmental Geochemistry Principle Geochemist and Co-Founder, Solana Beach, California

Education:

Ph.D., Geological Sciences (Geochemistry) - 1983; University of Southern California,

Los Angeles, California

M.S., Geological Sciences (Geochemistry) - 1978; University of Southern California,

Los Angeles, California

B.S., Chemical Engineering - 1975; Clarkson University,
Potsdam. New York

Summation of Employment History:

1990 to Date: Transglobal Environmental Geochemistry, Inc., Solana Beach, California,

Laboratory Director

1988 to Date: Transglobal Exploration and Geoscience, Inc., Solana Beach, California,

Vice President and Co-founder

1986 to 1988: InterOcean Systems, Inc., San Diego, California,

Chief Geochemist

1982 to 1986: Unocal Science and Technology Division, area, California

Research Geochemist

1977 to 1981: L.A. Pierce College, Los Angeles, California

Instructor

1977 to 1978: California State University Northridge, Northridge, California

Instructor

Professional Experience:

Dr. Hartman's daily responsibilities, with respect to TEG, include direction of laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review and method development, and direction of corporate development.

Dr. Hartman's academic research included a variety of chemical oceanography projects including: petroleum tracing using stable and radiogenic compounds as tracers; process controls across the airwater interface; and gas chromatography of dissolved gases in water and sediments.

Prior to co-founding Transglobal Exploration, Dr. Hartman was chief marine geochemist with InterOcean Systems, where his responsibilities included direction of marine sediment and bottom-water geochemical exploration programs. As a research chemist with Unocal Research, Dr. Hartman was responsible for development of geochemical prospecting techniques in onshore and offshore applications. These research operations included extensive method development and use of a variety of gas chromatography methods. Other research included use of stable metal isotopes as correlation parameters for organic materials.

Section 16 Revision 1 Date 1/1/93 Page 4 of 11

Vice President and Co-Founder:

Transglobal Exploration & Geoscience March 1988 to Present

Primary Duties:

As Co-founder and Vice President of Transglobal & Geoscience in 1988, Dr. Hartman has been personally responsible for directing geochemical technical development, equipment design and maintenance, applications development, and implementation of exploration programs, analyses, and reporting.

Marine Geochemist:

InterOcean Systems, Inc.
March 1986 to March 1988

Primary Duties:

Director of marine sediment and marine bottom-water geochemical exploration programs for petroleum.

Technical interpretation and report presentation of marine sediment and marine bottomwater (sniffing) hydrocarbon data fo. petroleum exploration.

Development of new analytical methods for marine sediment and marine bottom-water exploration for petroleum.

Research Geochemist:

Unocal Science and Technology Division September 1982 to March 1986

Primary Duties:

Director of two marine geochemical exploration programs for petroleum in the North Sea (1985) and the Bering Sea (1983).

Surface geochemical prospecting for precious metal deposits, petroleum, and geothermal resources.

Applications of radioisotopes for age dating petroleum and dating ground water residence times.

Investigations on the use of gaseous halocarbons as gas tracers for assessing the characteristics of petroleum reservoirs.

Pioneer studies on the applications of stable metal isotopes as correlation parameters for organic materials.

187309

Section 16 Revision 1 Date 1/1/93 Page 5 of 11

TEACHING

187310

Instructor:

Introductory Oceanography Laboratory, Physical Geology and Introductory Oceanography, L.A. Pierce College September 1977 to June 1981 (8 semesters)

Instructor:

Introductory Oceanography California State University Northridge September 1977 to May 1978

PUBLICATIONS

Publications in various geochemistry and marine chemistry fields. List available on request.

RESEARCH CRUISES

Over 40 oceanic geochemical research cruises.

DERHSING LUU, Ph.D.

187311

President, Transglobal Environmental Geochemistry/Texas

Education:

Ph.D., Analytical Chemistry

- 1986; Southern Illinois University,

Carbondale, Illinois

B.S., Chemistry

- 1977; National Chung Hsing University,

Taiwan

Summation of Employment History:

1992 to Date:

Transglobal Environmental Geochemistry/ Texas, Austin, Texas

President / Laboratory Director

1990 to 1991:

Texas Research Institute, Austin, Texas

Laboratory Manager, GC/MS Operator, Corporate QA Officer

1989 to 1990:

ENSR Corporation, Camarillo, California,

Air Toxics Monitoring Tech. Supervisor & Reg. QA Officer

1985 to 1989:

Global Geochemistry, Canoga Park, California

Project Manager

Protessional Experience:

Dr. Luu's daily responsibilities with respect to TEG/Texas include direction of laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review and method development.

Dr. Luu's academic research included a variety of novel automated analytical instruments development including: methanol analyzer and hydrocarbon analyzer.

Prior to founding TEG/Texas, Dr. Luu was R&D senior scientist with Global Geochemistry Corp. where his responsibility included development of methanol analyzer for CARB (California Air Resource Board) and hydrocarbon analyzer for Schlumberger. As a senior chemist for Air Toxic Lab with ENSR, Dr. Luu was responsible for daily laboratory operations and method development for air toxic laboratory. As an environmental laboratory manager and corporate QA officer with TRI (Texas Research Institute), Dr. Luu was responsible for supervising, performing GC/MS analysis, training, method developing for indoor air monitoring, and devising corporate QA/QC policy.

President and Director:

Transglobal Exploration & Geochemistry 1992 to Present

Primary Duties:

As President of TEG /Texas, Dr. Luu is in charge of direction of laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review and method development.

Section 16 Revision 1 Date 1/1/93 Page 7 of 11

Laboratory Manager, GC/MS Operator, Corporate QA Officer:

187312

Texas Research Institute 1990 to 1991

Primary Duties:

As Laboratory Manager, managed both EPA and IH work, method implementations, provided technical support for clients.

As GC/MS Operator, handled all GC/MS work including standard EPA methods (8240's, 624, 625, TO-14) and other non routine types of analyses such as material characterizations, indoor air analysis, etc.

As Corporate QA Officer, rewrote the corporate QA manual and laboratory QC handbook. Executed QA policy including internal audits, problem solving and provided QA knowledge for corrective actions.

Air Toxics Monitoring Technical Supervisor & Regional QA Officer: ENSR Corporation 1989 to 1990

Primary Duties:

Created the method development for air monitoring. Implemented the QA policy.

Project Manager:

Global Geochemistry 1985 to 1989

Primary Duties:

Project Manager for project entitled "Preconcentration and Determination of Reduced Sulfur Compounds in Ambient Air by Purge and Trap / GC-FPD Detection", project sponsored by Southern California Edison. Performed EPA analyses (EPA 8015, EPA 8020, and EPA 418.1). Designed a computer program for data calculations to improve EPA lab productivity and for instrument hardware control. R&D Project Manager for CARB research project to developed the methanol analyzer for exhaust emission studies.

MICHAEL A. KOROSEC, M.S.

187313

President, Transglobal Environmental Geosciences Northwest, Inc. Principle Geochemist, Environmental Chemist, Geologist Program Manager

Education:

MBA, Finance and Management

- 1987; City University and Pacific Lutheran Univ.

M.S., Geological Sciences

- 1978; University of Southern California

B.S., Biology

- 1975; Case Western Reserve University

Summation of Employment History:

1989 to Date:

Transglobal Environmental Geosciences Northwest, Inc.,

Laboratory Director and Analytical Chemist

1978 to 1989:

Washington Department of Natural Resources,

Geologist

1978 to 1980:

L.A. Pierce College

Instructor

1975 to 1978:

University of Southern California

Teaching Assistant and Instructor

1973 to 1975:

Chi Corporation
Technical Writer

Professional Experience:

Mr. Korosec's daily responsibilities with respect to TEG Northwest include direction of all laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review, marketing, and contract negotiations.

While with TEG, Korosec has developed unique preparation methods and analytical techniques for on-site analysis of PAH's, PCB's and chlorinated pesticides in soils, and PAH's and BTEX in tars.

Korosec's academic research included studies of the chemical and physical controls on the transport of dissolved nutrients across the sediment-water interface, using UV spectrophotometry and gas chromatography.

Prior to founding TEG/Northwest, Korosec was in charge of the state of Washington, Division of Geology and Earth Resources' Geothermal Exploration Program. The work included the development of a water analysis laboratory for the determination of dissolved cat ions, an ions and trace metals in thermal and mineral waters. Instrumentation included AA spectrophotometer, UV spectrophotometer, mercury analyzer and specific ion meters. Additional work included drilling programs for temperature gradient and heat-flow studies, whole rock geochemistry, age dating, and geologic mapping. As program manager, Korosec was responsible for all contracting, subcontracting and reporting to the U.S. Department of Energy.

187314

Laboratory Director and Analytical Chemist:

Transglobal Exploration & Geoscience 1989 to Present

Primary Duties:

Formed the Northwest branch of the TEG network through a contractual agreement with TEG, Solana Beach, CA. Own and operate a fleet of Mobile Environmental Laboratories.

Geologist:

Washington Department of Natural Resources: 1978 to 1989

Primary Duties:

State Geologic Map Program; production of 1:100,000 and 1:250,000 maps through compilation and new mapping. Program and Project Manager for U.S. Dept. of Energy-funded state-wide geothermal program. Geothermal exploration, construction of a complete water analysis laboratory, geochemical analysis of thermal and mineral waters, geothermal drilling.

Instructor:

L.A. Pierce College 1977 to 1978

Primary Duties:

Instructed course on Oceanography.

Teaching Assistant and Instructor:

University of Southern California 1975 to 1978

Primary Duties:

Instructed and assisted course on Oceanography.

Technical Writer:

Chi Corporation 1973 to 1975

Primary Duties:

Worked on newsletter, advertisements and user manual production.

Section 16 Revision 1 Date 1/1/93 Page 10 of 11

MARK JERPBAK, M.S., R.G.

187315

President, TEG Sacramento Director and Registered Geologist

Education:

M.S., Geology and Geophysics

- 1988; University of Iowa

Iowa City, Iowa

B.A., Geology

- 1981; Cal State University Chico

Chico, California

Summation of Employment History:

1992 to Date:

TEG Sacramento

Laboratory Director

1991 to Date:

Yolo County Environmental Health Services

Consultant

1990 to 1992:

EnLab Mobile Services

Marketing Director/Principal Analyst

1987 to 1990:

EnergyLog Corporation

Geologist/Geochemist

1984 to 1987:

University of Iowa

Graduate Student

(Research with Iowa Geological Survey)

1981 to 1984:

EnergyLog Corporation

Geologist/Geochemist

Professional Experience:

Mr. Jerpbak's daily responsibilities with respect to TEG/Sacramento includes direction of all laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review, marketing, and contract negotiations.

Since 1981, Mr. Jerpbak has worked with several companies and agencies in geology and geochemistry in the United States and overseas, including: Ascension Island, Papau New Guinea and Japan. His work has included gas chromatography in the petroleum industry, gas and vapor analyses and wet chemical titration in the geothermal industry, injection tests in active geothermal areas, and gas chromatography by EPA methods in the environmental industry. He has worked extensively in the field and in the mobile lab environment.

Mr. Jerpbak holds a M.S. degree from the University of Iowa in geology and geophysics. He is a California State Registered Geologist and also consults for the Yolo County Department of Public Health, and Environmental Health Services in California.

Laboratory Director:

TEG, Sacramento 1992 to Present

Section 16
Revision 1
Date 1/1/93
Page 11 of 11
187316

Primary Duties:

Mr. Jerpbak's daily responsibilities with respect to TEG/Sacramento includes direction of all laboratory operations, supervision of laboratory personnel, analysis, report preparation, technical review, marketing, and contract negotiations.

Consultant:

Yolo County Environmental Health Services 1991 to Present

Primary Duties:

Oversee continuing implementation of county ordinances regarding Class 2 injection wells. Geologic advisor to the Environmental Health Services Director.

Marketing Director / Principal Analyst:

EnLab Mobile Services
1990 to 1992

Primary Duties:

Directed Marketing and worked as principal analyst performing gas chromatography using EPA methods for organic analysis. A listed the building of a mobile laboratory.

Geologist:

EnergyLog Corporation

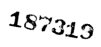
1981 to 1990 (Time off for University of Iowa 1984 to 1987)

Primary Duties:

Performed mobile lab work in the petroleum and geothermal industries located in California and overseas.

TAB

APPENDIX C



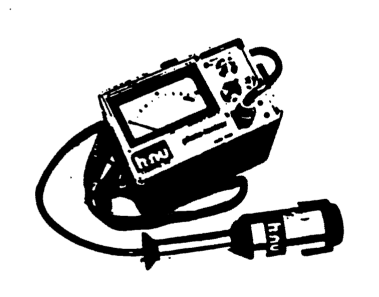


INSTRUCTION MANUAL

POR

MODEL PI 101

PHOTOIONIZATION ANALYZER



© Copyright 1878

TABLE OF CONTENTS

Section		Page
1	INTRODUCTION	1
2	OPERATION	5
3	CALIBRATION	19
4	DETECTION PRINCIPLE AND THEORY	22
5	TROUBLESHOOTING	45
6	REPLACEMENT PARTS LIST	55
7	MODEL PI-101 PHOTOIONIZATION ACCESSORIES	56
8	PI-101 RECORDER ASSEMBLY DISTRUCTIONS	57
9	WARRANTY	58

SECTION 1

INTRODUCTION

The model PI 101 has been designed to measure the concentration of trace gases in many industrial or plant atmospheres. The analyzer employs the principle of photoionization for detection. This process is termed photoionization since the absorption of ultraviolet light (a photon) by a molecule leads to ionization via:

where RH = trace gas

hy = a photon with an energy 2 Ionization Potential of RH

The sensor consists of a scaled ultraviolet light source that emits photons which are energetic enough to ionize many trace species (perticularly organics) but do not ionize the major components of air such as 0_2 , N_2 , ∞ , ∞_2 , or N_2 . A chamber adjacent to the ultraviolet source contains a pair of electrodes. Whe a positive potential is applied to one electrode, the field created drives any ions, formed by absorption of UV light, to the collector electrode where the current (proportional to concentration) is measured.

To minimize adsorption of various sample gases, the ion chamber is made of an inert fluorocarbon material, is located at the sampling point, and a rapid flow of sample gas is maintained through the small ion chamber volume.

The analyzer will operate either from a rechargeable battery for more than 10 hours or continuously from the AC battery charger. A solid state amplifier board in the probe and a removable power supply board in the readout module enabrapid servicing of the unit in the field.

The useful range of the instrument is from a fraction of a ppm to about 2,000 ppm. For measurement at levels above 2,000 ppm, dilution of the sample stream with clean air is recommended. Some typical specifications for the model PI 101 Photoionization Analyzer are given in Table 1.

TABLE 1

SPECIFICATIONS FOR MODEL PI 101 PHOTOIONIZATION ANALYZER

performance (benzene referred)

range 0.1 to 2000 ppm
detection limit 0.1 ppm:
sensitivity (mex) 0-2 ppm FSD over 100 division meter scale
repeatability + 1% of FSD
linear range 0.1 to 600 ppm
useful range 0.1 to 2000 ppm
response time < 3 sec to 90% of full scale
ambient hamidity to 95% RH
therating temperature ambient to 40°C*

physical

size: probe 6.3 DIA x 28.5L (cm) (2-1/2 x 11-1/4")
readout 21W x 13D x 16.5H (cm) (8-1/4 x 5-3/16 x 6-1/2
stored 21W x 13D x 24H (cm) (8-1/4 x 5-3/16 x 9-1/2
cable 80 cm long (32")

weight: probe .55 kg (20 cunces)

readour 3.2 kg (7 pounds)

total (shipping) 5.4 kg (12 pounds)

controls and functions

mode switch Off, Battery Check, Standby (zero), 0-2000, 0-200, 0-20 ppm low battery indicator light zero (10 turn + 300% FSD max) spen (10 turn counting dial 1.0 to 10 times nominal sensitivity) readour 4-1/2" (11.3 cm) meter Taux Bend movement graduated 0-5-10-15-20, divisions signal output for recorder 0-(-5V) FSD power output for recorder 12 VDC - jack on side of instrument

power requirements of operating times

continuous use, battery > 10 hours
continuous use with HNU recorder reduces instrument battery operating time
to 1/2 normal time
recharge time, max'< 14 hours, 3 hours to 90% of full charge
recharge current, max .4 Amps @ 15 VDC

TABLE 1 (Continued)

construction

Designed to withstand the shock and abuse to which portable instruments are often subjected. The readout is housed in a two piece aluminum case, and finished with a solvent resistant baked acrylic textured paint.

The probe is fabricated from extruded aluminum sections and machined plastic.

serviceability

The probe and readout are of a modular design allowing rapid servicing and/or replacement of machanical and electrical components. All module interwiring includes quick disconnects.

maintenance

The instrument contains only one moving part, and consumes no gases or reagents. The only routine maintenance procedure is cleaning the light source window every several weeks.

calibration check

Check instrument calibration at least once per week with HNU calibration standard to ensure that the high sensitivity of the instrument is maintained.

* Instrument is temperature compensated so that a 20°C change in temperature corresponds to a change in reading of < ± 27, full scale at maximum sensitivity.

SECTION 2

OPERATION

Umpack the instrument carefully and remove the housing, the probe and any 2.1 spare parts from the shipping carton. Place the instrument on a table or benc with the label upright. Remove the top section of the instrument by opening t two fasteners on the cover (see figure 1). The inner penel of the top section can be removed by pulling up on the fasteners. The top section of the instrument contains the battery charger and a waist strap. The waist strap clips on to the strap brackets of the instrument when needed.

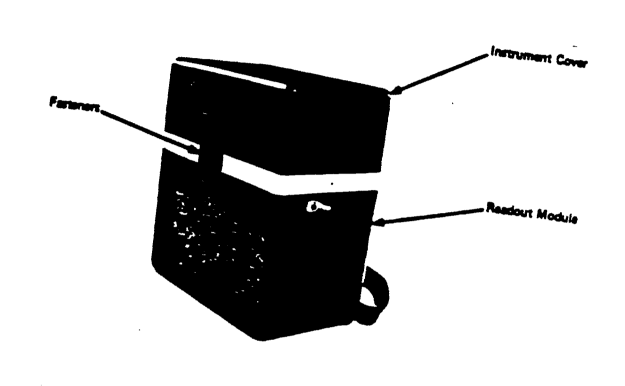
Before attaching the probe, check the function switch on the control pane to make sure it is in the off position. The 12 pin interface connector for th probe is located just below the span adjustment on the face of the instrument (see figure 2). Carefully match the Alignment Key in the probe connector to the 12 pin connector on the control penel, and then twist the probe corrector until a distinct snap and lock is falt.

Arrached to the insurement is a warranty card which should be filled our completely and returned to HNU Systems.

Then the function switch to the bettery check position. The needle on the 2.2 meter should read within or above the green battery arc on the scaleplate. It the needle is in the lower portion of the battery arc, the instrument should ! recharged prior to making any measurements. If red LED comes on, the battery should be recharged.

Next, turn the function switch to the on position. In this position the UV light source should be on. Look into the end of the probe to see the purple glow of the lamp.

A brief description of the instrument controls and functions is shown in Figure 2.



report 1. Unpacking the Photological

TABLE II

BRIEF DESCRIPTION OF DISTRIMENT CONTROLS AND FUNCTIONS*

Control

Function

Six Position Switch

OFF - Shuts off all power and removes DC vo

ON - In any other function position or the mode, the electronics are on.

BATTERY CHECK - Indicates the condition of battery. If needle position is in low portion of green battery art, the instrumnt should be recharged.

STANDBY - UV lamp is off but electronics are This position will conserve power and the useful operating time between right of the bettery. This position is a so lized to adjust the electronic zero.

RANCES - 0-20, 0-200, 0-2000 direct reseling evailable at minimum prin for benze e. sensitivity is available by adjusting span potentioneter.

Zero Potentiameter

ſ

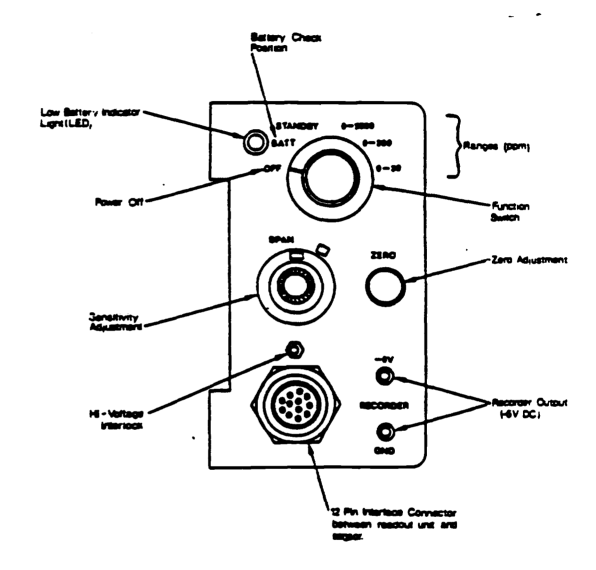
A ten turn potentiometer is employed to _; the zero electronically when the instrument placed in the standby position with the proactached. This eliminates the need for _; carbon free gas.

Span Potenticmeter

A ten turn counting potentioneter is utilize upscale setting of the meter on calibrator Counter-clockwise rotation increases the setting (~10 times). This pot can increase the sitivity to make the instrument direct reached any gas which the instrument rest reconstrument rest reconstrument.

*For position of layout controls see Figure 2.

Figure 2 Control Panel Functions



To zero the instrument, turn the function switch to the standby position and rotate the zero potentiometer until the meter reads zero. Clockwise rotation of the zero potentiometer produces an upscale deflection while countered the wise rotation yields a downscale deflection. Note: no zero gas is needed, since this is an electronic zero adjustment (see below). If the span adjustment set is changed after the zero is set, the zero should be reclacked and adjusted. If necessary, Wait 15 or 20 seconds to ensure that the zero reading is stable. If necessary, readjust the zero.

The instrument is now ready for calibration or measurement by switching the function switch to the proper measurement range. The instrument is suppled calibrated to read directly in ppm (v/v) 0-20, 0-200, 0-2000 of benzers with the span position set at 9.8. For additional sensitivity, the span potention to is turned counterclockwise (smaller numbers) to increase the gain. By changing the span setting from 10.0 to '0 the sensitivity is increased approximately and fold. Then, the 0-20, 0-200, and 0-2000 ppm scales become 0-2, 0-20, and 0-200 ppm full scale, respectively. This span control is also utilized to make the instrument scale read directly in ppm of the compound being measured. E.g., 't is adjusted to match the value of a calibration gas to that same reading on the instrument scale. The span control can be utilized to calibrate nearly any or appound, measured by photoionization, to be direct reading on the 0-20 ppm range. For example, gain settings of 4.5 or 8.9, respectively, will provide directly accapability (0-20, 0-200 ppm) for vinyl chloride and trichloroethylene, respectively.

A small DC operated fan is used to pull air through the photoionization sensor at a flow rate of three to seven handred contineters per minute (cs. 0.5 lpm). The fan provides nearly instantaneous response times (Figure 3) while consuming little power. The characteristics of a fan are such that it cannot tolerate a significant pressure drop without affecting the flow rate and therefore either the instrument reading or response time. Since photoionization is essentially a nondestructive technique, changes in flow rate do not affect the signal but if a large pressure drop is imposed at the inlet of the probe, the sample may not reach the sensor.

TABLE III

VERIFICATION OF ELECTRONIC ZERO FOR PHOTOIONIZATION ANALYZER*

Sample	Instrument Reeding (ppm)	% of F.S.
Room Air		4 01 P.S.
NOOM AIF	0.7	30
Room Air Passed Through	_	35
6" x 3/4" OD Charcoal	0.1	5
Scrubber		•
Zero Air		
	0.25	12.5
Zero Air Passed Through		12,3
6" x 3/4" OD Charcoal	0.04	2
Scrubber		_

^{*}Maximum Gain = 2 ppm full scale.

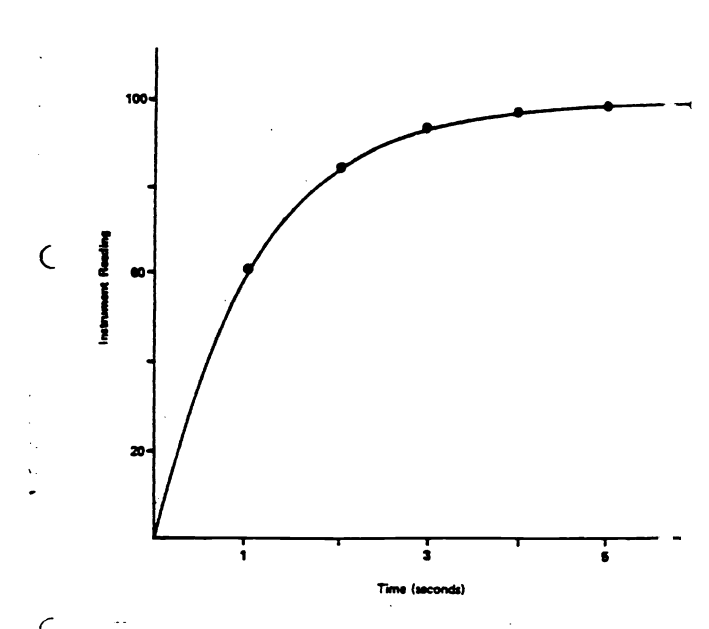
TABLE IV

RELATIVE PHOTOIONIZATION SENSITIVITIES* FOR VARIOUS GASES

Chemical Grouping	Relative Sensitivity	Examples
Arometic	10.0	Benzene, Toluene, Styrene
Aliphetic Amine	10.0	Diethytemine
Chlorinated Unsaturated	5-9	Vinyt Chloride, Vinvlidene Chloride, Trichloroethylene
Carbonyl	5- 7	MEK, MIBK, Acetone, Cyclohexene
Unsaturated	3-6	Acrolein, Propylene, Cyclohexene, Allyl Alcohol
Sulfide	3-5	Hydrogen Sulfide, Methyl Mercaptan
Paraffin (Cg-C7)	1-3	Pentane, Hexane, Heptane
Ammonia	0.3	•
Paraffin (C ₁ -C ₄)	0	Methane, Ethane

^{*}Sensitivities in ppm (v/v).

Figure 3. Time Response for the Photoionization Analyzer.



The instrument was designed to measure trace gases over a concentration range from less than 1 ppm to 2000 ppm. Higher levels of various gases (to percentage range) can be measured but the recommended procedure is to dilute the sample with clean air to a concentration of less than 500 ppm. This is generally within the linear range of the instrument and if the measured concentration is multiplied by the dilution ratio the correct concentration in the stream can be determined. A typical calibration curve is shown in Figure 4. Note that the calibration curve for benzers (the photoionization standard) is linear (over more than three decades) up to about 600 ppm (v/v).

If the probe is held close to AC power lines or power transformers, an error may be observed. For measurements made in close proximity to such items. their effect on measurements can be determined by the following procedure. Zero the instrument in an electrically quiet area, in the standby position, then move the instrument to the questionable area involved. If AC pickup is going to be a problem, the mater (in the standby position) will indicate the magnitude of

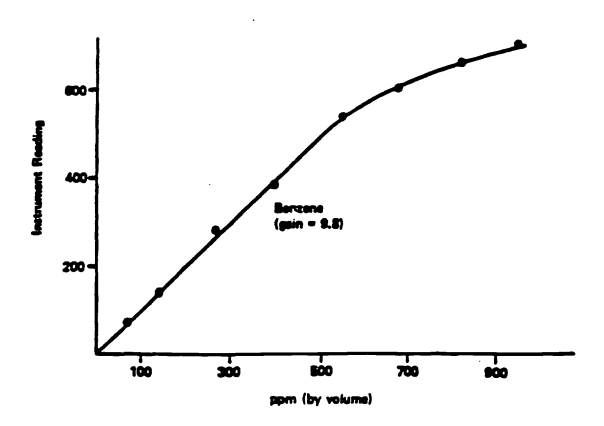
The instrument is equipped with an automatic solid state bettery protection circuit. When the bettery voltage drops below + 11 volts, this circuit will enconstically turn off the power to the instrument. This prevents deep discharging of the battery and considerably extends the battery life. If the instrument is unintentionally left on overnight, the bettery will be unharmed because of the bettery protection circuit. If the instrument bettery check reads low and the lamp doesn't fire, plug the charger into the instrument. The power to the enalyzer should then be returned.

To charge the battery, place the mini phone plug into the jack on left side of the bezel prior to plugging charger into 120 VAC. When discornecting charger, remove from 120 VAC before removing mini phone plug. The battery is completely recharged overnight (ca. 14 hours). To ensure that the charger is functioning, turn the function switch to the battery check position, place phosphug into jack and plug charger into AC outlet. The meter should go upscale if charger is working and is correctly inserted into the jack.

(

The instrument can be operated during the recharge cycle. This will lengthen the time required to completely recharge the instrument battery.

Figure 4. Typical Calibration Curve for Photoionization Analyzer.



CALLBRATION

Static or dynamic gas generation systems can be utilized for calibratic of the instrument. A number of such systems for generating test atmospheres for various gases have been described by G. O. Nelson in "Controlled Test Atmospheres," Arm Arbor Science Publishers, Arm Arbor, Michigan (1971).

The most convenient packages for calibration are the non-toxic analyzed gas mixtures available from HNU Systems in pressurized containers (Catalogue \$101-350).

A rapid procedure for calibration involves bringing the probe and readour in close proximity to the calibration gas, cracking the valve on the tank and checking the instrument reading. This provides a useful spot check for the instrument.

ment from a pressurized container is to connect one side of a "T" to the pressur container of calibration gas, another side of the "T" to a rotaneter and the side of the "T" directly to the 8" extension to the photoionization probe (see Figure 5). Crack the valve of the pressurized container until a slight flow s indicated on the rotaneter. The instrument draws in the volume of sample requir for detection, and the flow in the rotaneter indicates an excess of sample. Or adjust the span pot so that the instrument is reading the exact value of the call bration gas. (If the instrument span setting is changed, the instrument should be turned back to the standby position and the electronic zero should be reading if-necessary.)

T

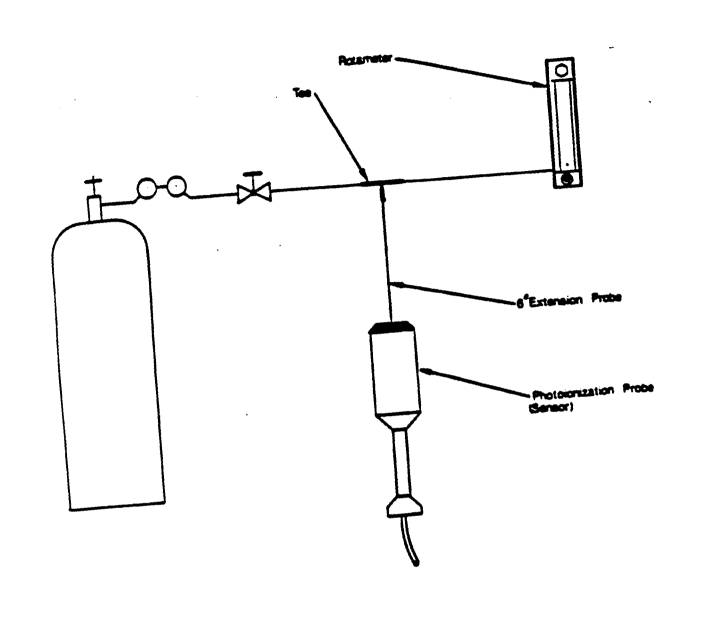


Figure S. Recommended Calibration Procedure for Photoionization Analyzar

The calibration gas should be prepared in the same matrix (air, nir n hydrogen, etc.) in which it is to be measured, otherwise an inaccurate read to may be obtained. The increased response which is seen in oxygen free gases can be attributed to a reduction in the quenching of ions by oxygen (actually () and is typical of any ionization detector. The quenching effect of oxygen is constant from about ten percent 0, to very high levels.

If a gas standard prepared in mitrogen is to be used for measurements in air, fill a 0.5 or 1 liter bag with the standard them add 50 or 100 cc of p recover to bring the level to 10-12%.

Any error between this value and 20% oxygen is quite small.

If the sample to be measured is in nitrogen, standards should be prepared in nitrogen. This will result in an increase in sensitivity of approximate.;

4.

^{*} Calibration with toxic gases should be performed in a hood since the 101 is a non-destructive analyzer.

SECTION 4

DETECTION PRINCIPLE AND THEORY

The detection principle of this portable instrument is photoionization. A wide variety of organic compounds and some inorganic compounds can be measured with this technique. Photoionization (with ~ 10 eV photons) applied to the analysis of trace gases in air can eliminate fragment ion formation (signals) from the major components of air yet still allow the ionization of many impurities of interest in industrial atmospheres. This is demonstrated by the listing of ionization potentials* in Tables V-XVII. Note the high (12 eV) ionization potentials for the major components of air. In addition, the choice of a sufficiently low ionization energy often permits the selective ionization of one or two components in a complex gas mixture.

While the ionization potential serves as a rough guide to whether or not a response is obtained, it does not predict what the quantitive response actually is. In some cases, a species with an ionization potential 10.3 or 10.4 eV will give a response. In these cases, however, the response is usually low because of its low ionization efficiency at 10 eV. A partial list of actual relative sensitivities obtained with a photoionization analyzer is given in Table XVIII. The use of the tables should allow a determination of the specificity of the instrument in a given application on many industrial processes; this instrument may not respond to the starting materials or by-products but will respond to a product. An example of this is seen in the vinyl chloride monomer plants where neither ethylene or dichloroethene is detected but vinyl chloride is detected.

^{*} Ionization potential is defined as the energy required to move an electron an infinite distance from the nucleus or more simply, the energy required to produce a positive ion and an electron.

READOUT UNIT

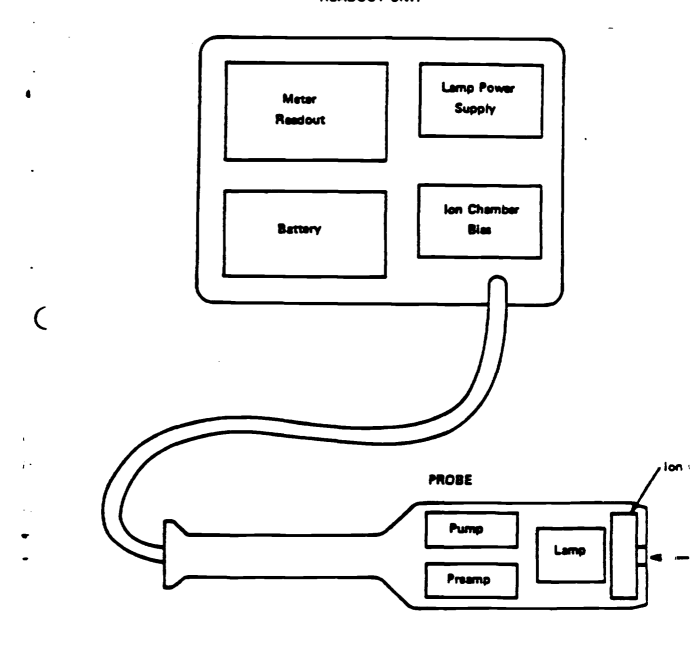


Figure & Block Diagram of Portable Photoionization Analyzer.

A block discrete of the major components of the Protoconics of the Pro is shown in Pigure 6. The instrument is separated into the writer intercentation Wellticordicur electrical cable. The Probe contains a few for source ind the season, the ultraviolet less which is ignited by applying a K solice. the state and carrows, the invitation character which contains a pair of The Process and is subjected to the lamp, and a stered amplified. The Process (4. 10 eV) which are and cred from the lamp pass through a W transmitting window and into the industrian distance absorption of the IV radiation by a solvening which has an ionization potential of 10 eV or less will lead to ion from the A Rosinively bissed high volcage electrode is used to Rush any inte Transaction of IV to the collector electronic later the contract. This corrected to 8 pro-PARTICISE BY the Expliciture in the Probe. In the Probe. Instrument is depicted in Flore 7. Fredrick and Provides & Politics and Politics and Polit The secretary of the inscrement is con-Tolled by charging the loop gain of the septimizer. A 12 volt bettery provides the Princes Princes for a light efficiency provides the provides of the princes. Porencials required for instrument operation.

Figure 7. Electrical Block Diagram of Photoconization analyze 187344 -4.5V Exha. Fan + 20V kon Chamber Gas Signal Lamp --450 PROBE Gain Control **€** 20 Recorper Output 27 Common DC-DC Conveter - - 10 VDC 19 Fan Voltage (OC) 7 21 - 16 VDC 12 V

READOUT ASSEMBLY

TABLE V

187345

SOME ATOMS AND SIMPLE MOLECULES

	IP (eV)		IP (eV)
Н	13.505	12	9.28
C	11.264	HF	15.77
N	14.54	HCI	12.74
0	13.614	HBr	11.62
Si	8.149	HI	10.38
S	10.357	so ₂	12.34
F	17.42	co ₂	13.79
CI	13.01	cos	11.18
Br	11.84	cs ₂	10.08
t	10.48	N ₂ O	12.90
H ₂	15.426	NO ₂	9.78
N ₂	15.580	03	12.80
02	12.075	H ₂ O	12.59
CO	14.01	H ₂ S	10.46
CN	15.13	H ₂ Se	9.88
NO	9.25	H ₂ Te	9.14
СН	11.1	HCN	13.91
ОН	12.18	C ₂ N ₂	13.8
F ₂	15.7	NH ₃	10.15
Cl ₂	11.48	CH ₃	9.840
Br ₂	10.55	CH ₄	12.98

TABLE VI

PARAFFINS AND CYCLOPARAFFINS

Molecule	IP (eV)
methane	12.98
ethane	11.65
propane	11.07
n-butane	10.63
i-butane	10.57
n-pentane	10.35
i-pentane	10.32
2,2-dimethylpropane	10.35
n-hexane	10.18
2-methylpentane	10.12
3-methylpentane	10.08
2,2-dimethy/butane	10.06
2,3-dimethy/butane	10.02
n-heptane	10.08
2.2,4 trimethylpentane	9.86
Cyclopropane	10.06
Cyclopentane	10.53
Cyclohexane	9.88
methylevelohevane	6.05

TABLE VII

187347

	ALKYL HALIDES	187347
Molecule		₽ (eV)
HCI		12.74
CI ₂		11.48
CH ₄		12.98
methyl chloride		11.28
dichloromethene		11.35
trichloromethane		11.42
tetrachioromethane		11.47
ethyl chloride		10.98
1,2-dichloroethane		11.12
1-chloropropane		10.82
2-chloropropane	•	10.78
1,2-dichloropropane	· · · · · — · · ·	10.87
1,3-dichloropropane		10.85
1-chlorobutane		10.67
2-chlorobutane		10.65
1-chloro-2-methylpropane		10.66
2-chloro-2-methylpropane		10.61
HBr		11.62
Br ₂		10.55
methyl bromide		10.53
dibromomethene		10.49
Tibromomethane		10.51
CH ₂ 8rCl		10.77
CHBr ₂ CI		10.59
ethyl bromide		10.29
1,1-dibromoethane		. 10.19
1-bromo-2-chloroethane		10.63

TABLE VII (Continued)

Molecule	IP (eV)
1-bromopropane	10.18
2-bromopropane	10.075
1,3-dibromopropane	10.07
1-bromobutane	10.13
2-bromobutane	9.98
1-bromo-2-methylpropane	10.09
2-bromo-2-methylpropane	9.89
1-bromopentane	10.10
н	10.38
12	9.28
methyl iodide	9.54
diiodomethane	9.34
ethyl iodide	9.33
1-iodopropane	9.26
2-iodopropene	9.17
1-iodobutane	9.21
2-iodobutane	9.09
1-iodo-2-methylpropene	9.18
2-iodo-2-methylpropane	9.02
1-iodopentane	9.19
F ₂	15.7
HF	15.77
CFCI ₃ (Freon 11)	11.77
CF ₂ Cl ₂ (Freon 12)	12.31
CF ₃ CI (Freon 13)	12.91
CHCIF ₂ (Freon 22)	12.45
CEBra	10.67

187349

TABLE VII (Continued)

Molecule	₽ (eV)-
CF ₂ 8r ₂	11.07
CH3CF2CI (Genetron 101)	11.98
CFCI ₂ CF ₂ CI	11.93
CF ₃ CCl ₃ (Freon 113)	د11.7 د
CFH8rCH ₂ 8r	10.75
CF ₂ 8rCH ₂ 8r	10.83
CF3CH2I	10.00
n-C ₃ F ₇ I	10.36
n-C ₃ F ₇ CH ₂ Cl	11.84
n-C ₃ F ₇ CH ₂ I	9.96

TABLE VIII

ALIPHATIC ALCOHOL, ETHER, THIOL, AND SULFIDES

Molecule	IP (eV)
H ₂ O	12.59
methyl alcohol	10.85
ethyl alcohol	10.48
n-propyi alcohol	10.20
i-propyl alcohol	10.16
n-butyl alcohol	10.04
dimethyl ether	10.00
diethyl ether	9.53
n-propyl ether	9.27
i-propyl ether	9.20
	10,46
H ₂ S	
methanethiol	9.440
ethanethiol	9.285
1-propanethiol	9.195
1-butanethiol	9.14
dimethyl sulfide	8.685
ethyl methyl sulfide	8.55
diethyl sulfide	8.430
dispersive sulfide	8.30

TABLE IX

ALIPHATIC ALDEHYDES AND KETONES

Malecule	₽ (eV)
co ₂	13.79
formaldehyde	10.87
acetaldehyde	10.21
propionaldehyde	9.98
n-butyraldehyde	9.86
leobutyraidehyde	9.74
n-valeraldehyde	9.82
isovaleraldehyde	9.71
acrolein	10.10
crotonaldehyde	9.73
benzaldehyde	9,53
ecetone	9.69
methyl ethyl ketone	9.53
methyl n-propyl ketone	9.39
methyl i-propyl ketone	9.32
diethyl ketone	9.32
methyl n-butyl ketone	9.34
methyl l-butyl ketone	9.30
3,3-dimethyl butanone	9.17
2-heptanone	9.33
cyclopentanone	9.28
cyclohexanone	9.14
2,3-butsnedions	9.23
2,4-pentanedione	8.87

TABLE X

ALIPHATIC ACIDS AND ESTERS

Molecule	IP (eV)
co ₂	13.79
formic scid	11.05
scetic soid	10.37
propionic acid	10.24
n-butyric scid	10.16
isobutyric scid	10.02
n-valeric acid	10.12
methyl formate	10.815
ethyl formate	10.61
n-propyl formate	10.54
n-butyl formate	10.50
isobutyl formate	10.46
methyl acetate	10.27
ethyl acetate	10.11
n-propyl acetate	10.04
isopropyl acetate	9.99
n-butyl acetate	10.01
isobutyl acetate	9.97
sec-butyl acetate	9.91
methyl propionata	10.15
ethyl propionata	10.00
methyl n-butyrate	10.07
methyl isobutyrate	9.98

TABLE XI

187353

ALIPHATIC AMINES AND AMIDES

Molecule	₽ (eV)
NH ₃	10.15
methyl amine	8.97
ethyl amine	8.86
n-propyl amine	8.78
i-propyl amine	8.72
n-butyl amine	8.71
i-butyl amine	8.70
s-butyl amine	8.70
t-butyl amine	8.64
dimethyl amine	8.24
diethyl amine	8.01
di-n-propyl amine	7.84
di-i-prop , , amine	7.73
di-n-butyl amine	7.69
trimethyl amine	7.82
triethyl amine	7.50
tri-n-propyl amine	7.23
formemide	10.25
acetamide	9.77
N-methyl acetamide	8.90
N,N-dimethyl formsmide	9.12
N,N-dimethyl acetamide	8.81
N,N-diethyl formamide	1.89
N.N-diethyl acetamide	8.60

187354

TABLE XII

OTHER ALIPHATIC MOLECULES WITH N ATOM

Molecule	IP (eV)
nitromethane	11.08
nitroethane	10.88
1-nitropropane	10.81
2-nitropropane	10.71
HCN	13.91
acetonitrile	12.22
propionitrile	11.84
n-butyronitrile	11.67
acrylonitrile	10.91
3-butene-nitrile	10.39
ethyl nitrate	11.22
n-propyl nitrate	
methyl thiocyanata	10.065
ethyl thiocyanate	9.89
methyl isothiocyanate	9.25
ethyl isothiocyanate	9.14

0-37

TABLE XIII

OLEFINS, CYCLO-OLEFINS, POLENES, ACETYLENES

Molecule	P (eV)
ethylene	10.515
propylene	9.73
1-butene	9.58
2-methylpropene	9.23
trans-2-buttene	9.13
cis-2-buttens	9.13
1-pentene	9.50
2-methyl-1-buttene	9.12
3-methyl-1-butane	9.51
3-methyl-2-butane	8.67
1-hexene	9.46
1,3-butadiene	9.07
Isoprene	8.845
cyclopentane	9.01
cyclohexene	2.945
4-methylcyclohexene	8.91
4-cinylcyclohexene	8.93
cyclo-ocustratene	7.90
acetylene	11.41
ргорупе	10.38
1-butone	10.12

TABLE XIV

SOME DERIVATIVES OF OLEFINS

Molecule	IP (eV)
vinyl chloride	9.995
cis-dichloroethylene	9.65
trans-dichlorosthylene	9.66
trichloroethylene	9.45
tetrachloroethylene	9.32
viny! bromide	9.80
1,2-dibromoethylene	9.45
tribromoethylene	9.27
3-chloropropene	10.04
2,3-dichloropropene	9.82
1-bromopropene	9.30
3-bromopropene	9.7
CF3CCI=CCICF3	10.38
n-C ₅ F ₁₁ CF=CF ₂	10.48
acrolein	10.10
crotonaldehyde	9.73
mesityl oxide	9.08
vinyl methyl ether	8.93
allyl alcohol	9.67
vinyl acetate	9.19

TABLE XV

HETEROCYCLIC MOLECULES

Molecule	IP (eV)
furan	8.89
2-methyl furan	0.39
2-fursidehyde	9.21
tetrshydrofuren	9.54
dihydropyran	8.34
tetrahydropyran	9.26
thiophene	8.860
2-chlorothiophene	8.68
2-bromothiophene	1.63
руттоїе	8.20
pyridine	9.32
2-picoline	9.02
3-picoline	9.04
4-picoline	9.04
2,3-lutidine	8.85
2.4-lutidine	8.85
2.6-juridine	8.85

TABLE XVI

AROMATIC COMPOUNDS

Molecule	IP (eV)
benzene	9.245
toluene	8.82
ethyl benzene	8.76
n-propyl benzene	8.72
i-propyl benzene	8.69
n-butyl benzene	8.69
s-butyl benzene	8.68
t-butyl benzene	8.68
o-xylene	8.56
m-xylene	8.56
p-xylene	8.445
mesitylene	8.40
durane	8.025
styrene	8.47
a-methyl styrene	8.35
ethynylbenzene	8.815
nepthalene	8.12
1-methylnapthalene	. 7.96
2-methylnapthalene	7.955
biphenyl	8.27
phenol	8.50
anisole	8.22
phenetole	8.13
benzaldehyde	9.53
acetophenone	9.27
benzenethiot	8.33
phenyl isocyanate	8.77

187353

TABLE XVI (Continued)

Molecule		P (eV)
phenyl isothiocyanete		8.520
Senzonitrile		9.705
nitrobenzene		9.92
aniline		7.70
fluoro-benzene		9.195
chicro-benzene		9.07
bromo-benzene		8.98
iodo-benzene		8.73
o-dichlorobenzene		9.07
m-dichlorobenzene		9.12
p-dichloroberzene		8.94
1-chloro-2-fluorobenzene		9.155
1-chioro-3-fiuorobenzene		8.21
1-bromo-4-fluorobenzene	**	8.99
o-fluorotoluene		8.915
m-fluoratoluene		8.915
p-fluorotoluene		8.785
o-chlorotoluene		8.83
m-chlorotoluene		8.83
p-chlorotolusne		€.70
o-bromotoluene		8.79
m-bromotoluene		8.81
p-bramataluene		8.67
o-iodotoluene		8.62
m-iodataluene		8.61
p-iodataluene		8.50
benzotrifluoride		9.68
o-fluorophenol		8.66

TABLE XVII

MISCELLANEOUS MOLECULES

Molecule	iP (eV)
ethylene oxide	10.565
propylene oxide	10.22
p-dioxane	9.13
dimethoxymethane	10.00
diethoxymethane	9.70
1,1-dimethoxyethane	9.65
propiolectone	9.70
methyl disulfide	8.46
ethyl disulfide	8.27
diethyl sulfits	9.68
thiolacetic acid	10.00
acetyl chloride	11.02
acetyl bromide	10.55
cyclo-C ₆ H ₁₁ CF ₃	10.46
(n-C ₃ F ₇)(CH ₃)C=0	10.58
trichlorovinylsilane	10.79
(C ₂ F ₅) ₃ N	11.7
isoprene	9.08
nhorene	11.77

TABLE XVIII

RELATIVE SEISTITIVITIES FOR VARIOUS CASES (10.2 eV Lamp)

187361

Species	Photoionization Sensitivity*
p-xylene	11.4
n-xylene	11.2
benzene	10.0 (reference standard)
tolume	10.0
distiyl sulfide	10.0
disthyl snine	9.9
Styrens	9.7
trichloroethylene	8.9
carbon disulfide	7.1
isobutylene	7.0
acatone	6.3
tetrahydrofur.	6.0
methyl ethyl ketone	5.7
methyl isoburyl ketone	5.7
cycloheumone	5.1
napths (86% aromatics)	5.0
Vinyl chloride	5.0
methyl isocyanate	4.5
iodine	4.5
methyl mercaptan	4.3
directly sulfide	4.3
allyl alcohol	4.2

(Species	Photoionization Sensitivity*
	propylene	4.0
	mineral spirits	4.0
	2,3-dichloropropene	4.0
	cyclohexene	3.4
	crotonaldehyde	3.1
	acrolein	3.1
	pyridine	3.0
	hydrogen sulfide	2.8
	ethylene dibromide	2.7
	n-octane	2.5
_	acetaldehyde oxime	2.3
	hexane	2.2
	phosphine	2.0
	heptane	1.7
	allyl chloride (3-chloropropene) ethylene ethylene oxide	1.5 1.0 1.0
	acetic anhydride	1.0
	c pinene	0.7
	dibramchloropropæne	0.7
	_ epichlorohydrin	0.7
	nitric oxide	0.6
	s pineme	0.5
	citral	0.5
	amonia	0.3

Species	Photoionization Sensitivity#	
acetic acid	0.1	
nitrogen dicadde	0.02	
methane	0.0	
acet, lare	0.0	
ethylene	0.0	

^{*}Expressed in ppm (v/v).

TROUBLESHOOTERS

If problems occur while using the photoionization enalyzer, it is recommended that the following troubleshooting guide be followed before consulting the factory.

5.1 General Aid to Fault Determination

Check battery condition. Recharge if necessary (Section 2). Turn the instrument on. Look into the Sample Inlet of the probe unit. A violet colore glow from the UV light source should be observed in all positions of the mode switch except the standby position. If unstable readings are obtained a fault probe cable or electrical connection could be the problem. To check, hold the probe normally and flex the cable firmly. Watch the mater for fluctuations as the cable is stressed. Individual wires in the readout can be checked in a similar way. Check the coordal connector on the amplifier board in the probe.

In the more sensitive ranges, a fluctuation in the reading may be noted if a hand or other large object is placed in very close proximity to the prol. This is normal for the instrument and will not result in an error in the measurement as long as the probe is held stationary while the measurement is being to en

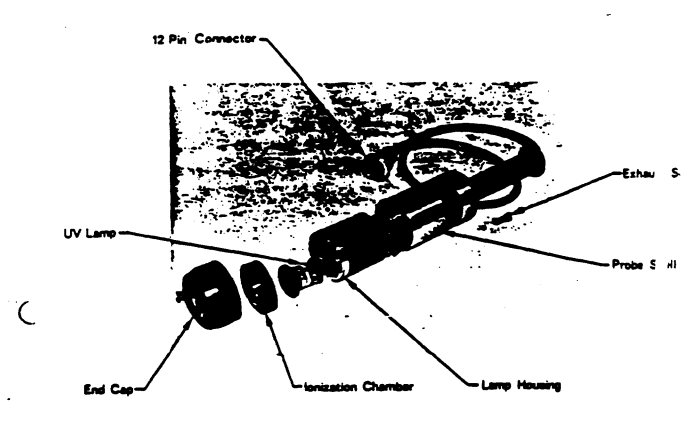
If the probe is held close to AC power lines or power transformers an error may be observed. If measurements are to be made in close proximity to such it is their effect on measurements can be determined by the following procedure. Zero the instrument in an electrically quiet area in the standby position, then move the instrument to the questionable area involved. If AC pickup is going to be a problem, the meter (in the standby position) will indicate the magnitude of the error.

5.2 Disassembly of Instrument

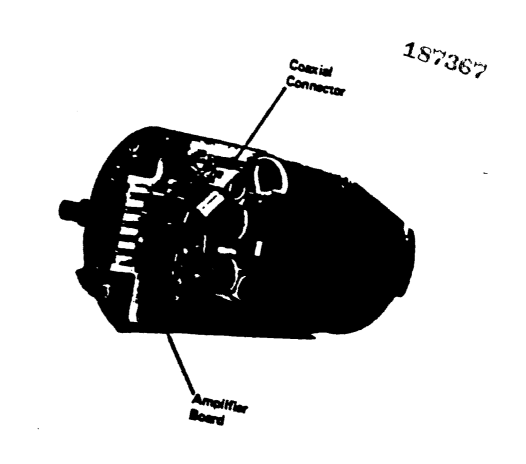
PROBE - Turn the function switch to the OFF position and discorrect the probe connector from the readout unit. Remove the enhant screw found near the base of the probe (see Figure 8.) Grasp the end cap in one hand and the probe shell in the other, and gently pull to separate the end cap and lamp housing from the shell. Loosen the screws on the top of the end cap and separate the end cap and ion chamber from the lamp and lamp housing. Gave must be taken so that the ion chamber doesn't fall out of the end cap and the lamp obesn't slide out of the lamp housing. Turn the end cap over in your hand and tap on the top of it; the ion chamber should fall out in your hand. Flace one hand over the top of the lamp housing and tilt slightly; the light source will slide our of the housing. The amplifier board can be removed from the lamp housing assembly by unmapping the control connection and then removing the retaining screw.

To reassemble this unit, first slide the lamp back into the lamp housing. Place the ion chamber on top of the lamp housing, checking to make sure that the contacts are properly aligned. Place the end cap on top of the ion chamber and replace the two screws. The screws should be tightened only enough to seal the "U" ring. DO NOT OVERTIGHEN. Line up the pins on the base of the lamp housing with the pins inside the probe shell. Gently slide the housing assembly into the probe shell. DO NOT FORCE the assembly into the shell as it only fits one way.

READULT UNIT - Turn the function switch to the CEF position and discorrect the probe from the readout unit before disassembly is conducted (see Figure 10). Remove the accessory power jack plug. Loosen the screw on the bottom of the case and, holding the instrument by the bezal, remove the case. The power supply board and control panel can be removed by unscrewing two screws and two muts. The entire panel, including the function switch, zero and span pots is removed in this operation Electrically discorrecting this module is simple, since all connections are made with Holex connectors.



Finance Community Parts of Probe



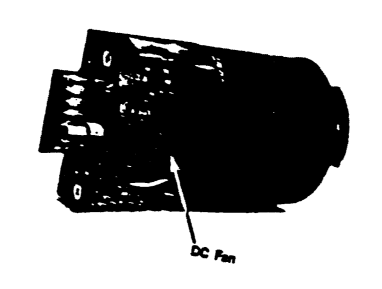


Figure 8. Component Parts of Lamp Housing.

5.3 Specific Faults

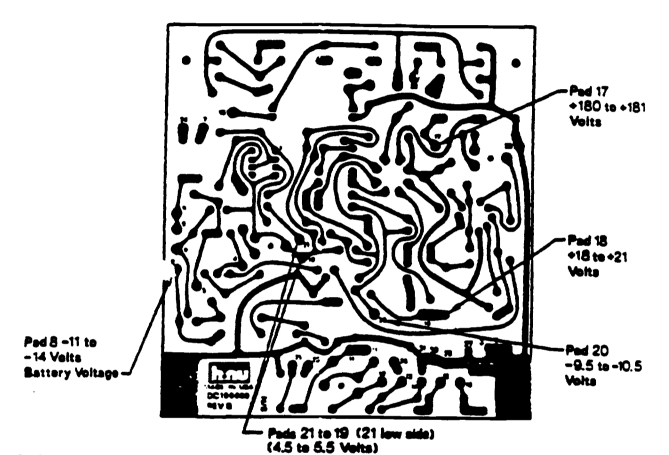
- 1. No meter response in any switch position (including BATT CHE)
 - A. Broken meter movement
 - (1) Tip instrument rapidly from side to side. Heter needle should move freely, and return to zero.
 - B. Electrical connection to meter is broken
 - (1) Check all wires leading to meter and clean the contacts of quick-disconnects.
 - C. Battery is completely dead
 - (1) Discorrect battery and check voltage with a volt-ohm meter.
 - D. Check 2 amp fuse

. (

11

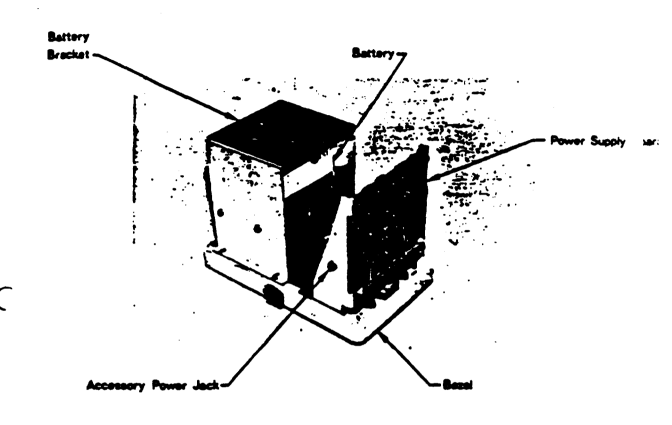
- E. If none of the above solves the problem, consult the factory.
- 2. Meter responds in BATT CHK position, but reads zero or near zero for all others
 - A. Power supply defective
 - (1) Check power supply voltages per Figure 11. If any voltage is our of specification, consult the factory.
 - B. Input transistor or amplifier has failed
 - (1) Rotate zero control; mater should deflect up/down as control is
 - (2) Open probe. Both transistors should be fully seated in sockets
 - C. Input signal connection broken in probe or readout
 - (1) Check input connector on printed circuit board. Should be firmly pressed down.
 - (2) Check components on back side of printed circuit board. All cornect: should be solid, and no wires should touch any other object.
 - (3) Check all wires in readour for solid connections.
- 3. Instrument responds correctly in BATT CHK, AND STBY, but not in measuring TOC
 - A. Check to see that light source is on (See General Faults section.)
 - (1) Check high voltage power supply (see Figure 11).
 - (2) Open and of probe, ramove lamp and check high voltage on lamp contact ring.
 - (3) If high voltage is present at all above points, light source has: st likely failed. Consult the factory.

- L All voltages measured with probe connected and instrument mode switch in BATT CHK position.
- All measurements referred to ground (pads 2.3 and 27) except pad 21 measured to pad 19 and pad 8 to pad 11.

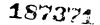


All Voltages Respect to Ground pede voltage pods veltage veltage voltage seds - 5.7V 1 9 - 12.2V 17 180V 25 0 2 GRD 10 - 12.1V 18 - 19AV 26 0 3 GRD 11 0 19 - 106V 27 GAD 4 -107V 12 0 20 - 97V 28 0 5 - 113V 13 Ō 21 - 14.57 GRD 29 6 - 12.1V 14 0 22 -400V 30 GRD 7 0 15 0 23 0 31 GRD 8 - 12.2V 16 ٥ 24 0

Figure 11 Power Supply PC Board



Shore St. Commoney Davis Of Seedard Unit.





To: Sales

From: Q.C.
Date: 12-21-94
Subject: Spare Probe(s) Calibration of PI-101 for

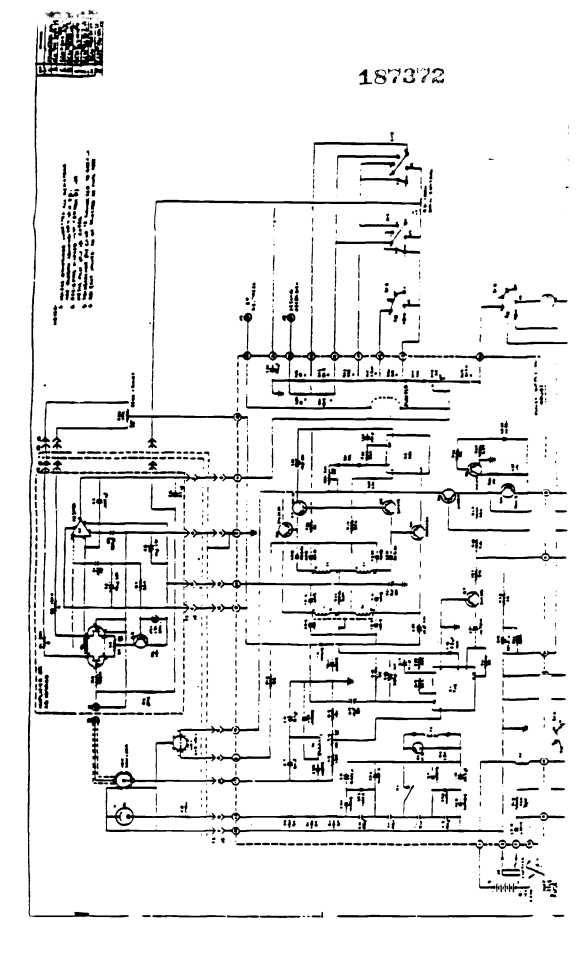
WO# 116. LAW ENGINEERING AND TESTING

A PI-101. S/N $\frac{41653}{1000}$, was calibrated to be direct reading on Benzene with two probe assemblies, consulting of a $\frac{10.2}{1000}$ eV lamp ($\frac{12696}{13387}$) probe (S/N $\frac{41653}{10000}$) and a $\frac{11.7}{10000}$ eV lamp ($\frac{13387}{10000}$) probe (0.4507) respectively. Calibration was performed as follows: The __IQ.2__ eV lamp probe assembly was connected to the unit and calibrated to be direct reading on benzens. The __II____ eV lamp probe assembly was then connected and calibrated by rezeroing the unit and adjusting the span to ______ to read benzene directly. Rezeroing must be performed with every probe assembly interchange.

The pertinent data is shown in Table 1. The unit is now calibrated to read benzene directly for both probe assemblies, with the same power supply module. It is important to emphasize that the isings cannot be interchanged between the two probes, aince the individual ampuriers are calibrated to lamp output.

Table 1

Power Supply S/N	Probe Amp S/N	Lamo (eV)	Lamp (S/N)	Span — Pot	100 ppm Benzene	HNU
41653	4051	10,2	12696	9,8	100.0	_5
41653	4160	_11,7_	13387	4.0	100.0	

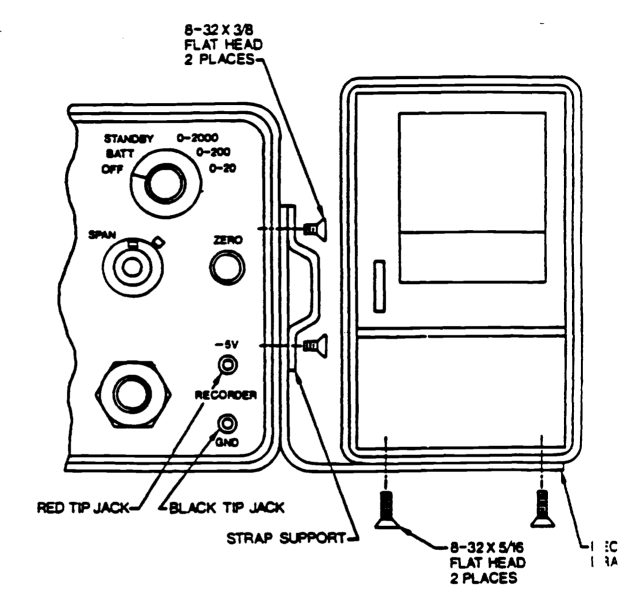


SECTION 6

REPLACEMENT PARTS LIST

187373

	Part No.	Assembly
	100004A	Probe Shell Assembly
	100005A	Ion Chamber
•	100006A	Fen/Light Source Housing
	100007A	Amplifier Board
	100008A	Fen
••	100009A	Light Source (10.2 eV)
	100010A	'O' Ring Kit
	100011A	Battery
	100012A	Yeter
	100013A	Charger
. (100014A	Power Supply Unit
	100015A & 100016A	Case (both halves)
•	100017A & 100018A	Straps
•	100019A	Switch
•	100020A	Pot (spen)
	4750-0001A	Pot (zero)
• :	100074	Probe Extension



NOTES:

- 1. Remove strap support on right side of FI-101 case.
- 2. Hunt recorder bracket between case & strap support using 8-32 X 3/8 long screws s pl
- 3. Hount recorder to bracket using 8-32 X 5/16 screws supplied.
- 4. Connect harness to plug on rear of recorder. Insert power plug into receptacle on side of case. Insert red pin plug into red tip jack and black pin plug into black tip jack on PI-101 front panel.

187375

Publications on Photoionization Available from HNU Systems

- (1) Driscoll, J. N. and P. Warneck, "The Analysis of ppm Levels of Gases in Air by Photoionization Mass Spectrometry," J. Air Poll. Cont. Assoc. 23, 858 (1973).
- (2) Driscoll, J. N. and F. F. Spaziani, "A New Instrument for Continuous Monitoring of Odorous Sulfur Compounds," presented at the ISA Mational Meeting, N.Y. (Oct. 1974).
- (3) Driscoll, J. N. and F. F. Spazieni, "Trace Gas Analysis by Photoionization" presented at the ISA Analysis Instrumentation Div. Meeting, King of Prussis, Ps. (May 1975).
- (4) Photoionization Detector for Gas Chromatography," presented at the Pittsburgh Conf. on Anal. Chem. and Spectroscopy, Cleveland (March 1976).

Requests for these papers should be sent to:

Publications Department

HNU SYSTEMS, INC. 160 CHARLE 'YT ST. NEWTON HIGHLANDS, MA. 02161

(617) 964-6690

Local Representative:

Tany Daniels (404) 991-7170

ADDENDUM A

to

Instruction Manual

for

Model PI 101

PHOTOIONIZATION ANALYZER

ION CHAMBER CLEANING PROCEDURE

Although the technique of photoionization is not sensitive to moisture, some electronic instability can be seen in the 101 portable analyzer as the result of excessive moisture. The following chart lists the symptoms expected, the possible causes, and solutions:

Symptom	Possible Cause	Solution
Loss in sensitivity	Condensation on instrument window	Avoid extreme temperature changer
		Acclimatize in- strument to environ- ment
Zero drift	Condensation on polarizing electrode	•
	Ion chamber is	Clean ion chamber

To clean the ion chamber: (See Section 5.2)

Remove the ion chamber. Remove all four screws, swing screen aside, and puriout the mesh. Using a Q-tip, gently clean the chamber with methanol to remove deposit. Dry in a vacuum oven at 90°C for 2 hours. Reassemble.

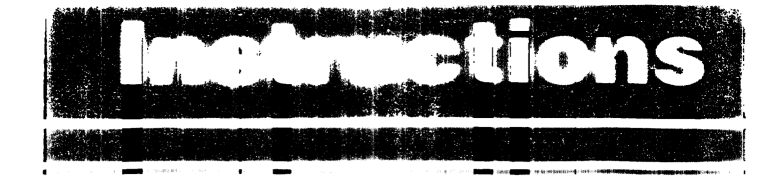
dirty

HNU MODEL 101 SPECIFICATION SHEET

	Serial # 4/6	<u>53</u>
Calibration BENZEN	E bySD_	
Range: 0-20, 0-200	7-2000 FFM	
Spen Pot Setting: 9,8	(10, ZEV) , 4,0 (11.7	€V)
	10.2 eV 12696 1	
Sensitivity:		
Detection Limit:		
Accessories:		
GAS	CONC (PPM)	RESPONSE
PROBE SN 41453 (10 BENZENE	CONC. (PPM) 100,0	100,0
HNU REF	97,0	<i>5</i> 6, 0
- 2	7)	
FR-08E SN 04507 (11.		100.0
BENZENE	100,0	100,0
HALL REF	97,0	79.0

TAB

APPENDIX D



Accided Mode 956 Portable Analog H/m/ Meter

(atalog No. 13-636-956







187382

CONTENTS

Title

PERFORMANCE CHARACTERISTICS	1
SPECIFICATIONS	1
INTRODUCTION	1
UNPACKING	2
OPERATING CONTF:OLS	
PRELIMINARY PROCEDURES	3
Checking and Adjusting Meter Zero	4
Installing pH Electrode System	4
OPERATION	4
Standardizing Instrument	5
Performing pH Determinations	5
Performing Millivott Determinations	6
SERVICE	6
Troubleshooting Chart	7
Checking Instrument Performance	8
Checking Buffer Accuracy	8
Checking Electrode System Efficiency	
Removing Meter Panel	8
Calibration Milliant Scale	8
Calibrating Millivolt Scale	9 10
Replacement Parts.	10
ILLUSTRATIONS	
-	.

re ·	Page
ntrois	3
rnal Components	9
nematic Diagram C69364	
se Assembly D69368	
ter Assembly C69363	13
Board Assembly R41840	14

Rende:

pН

0 - 14 pH

Millivolts

= 700 mV

Resolution: pН Milliwolte

0.05 pH 5 mV

Precision:

DH Millivolts ± 0.05 pH ±5 mV

Relative Accuracy:

= 0.05 pH'

SPECIFICATIONS

input impedance

Tolerable Source Resistance

Amplifier Drift Manual Temp. Compensation

Battery

Panel Meter

Environmental Rating:

Humidity Range Operating Temp. Range Storage Temp. Range

Physical Data: **Dimensions**

Carrying Weight Shipping Weight >2.5 x 1016 ohms

0-2000 megohms

0.5 pH in 24 hrs.

0 to 100°C

9 volt alkaline²

41/2" taut band with mirror-back

187383

0 - 90%

15 to 45°C 0° to 80°C

34.3cm W x 25.4 D x 10.2 H

(13.5W x 10 D x 4 H)

1.8 kg (4 lb.) 2.3 kg (5 lb.)

INTRODUCTION

The Fisher Model 956 Accumet pH meter is a portable, battery-powered instrument intended for field use where rough handling can be expected and where electric power is not available. Mounted on the front panel are three simple operating controls used to select measuring function (pH or mV), to standarr instrument with a buffer solution, and to compensate for sample temperature. The meter is equippe. BNC and pin input jacks. A threaded sycket accepts the electrode support rod. The meter movement and associated electronics are mounted on the rear of the panel, and can be completely removed from the Ci for service (and battery replacement) by simply removing four screws.

The Model 956 is powered by a supplied 9-volt alkaline battery with an operating life of approximate 10 hours. The instrument employs advanced operational amplifier input circuitry, thus using to advantage the latest in solid-state technology for optimum reliability and trouble-free service in the most demanding environments.

^{&#}x27;Within 2 pH units of standardization.

²Approximately 100 hrs. operating time.

UNPACKING

The Accumet Model 956 is shipped in a single carton containing the items listed below. While unbacking, check each item against the packing list. If a shortage exists, notify your representative, giving the name and number of the missing item. Also be sure to complete and return the warranty card located at the rear of this manual.

NOTE: In the event that shipping damage has been observed, retain the carton and packing material intact with the unit and file a claim with the final carrier. Usually, the firm will send an inspector to ascertain liability.

Quantity	item .	Cat. No. or Part No.
1	Accumet Portable pH Meter Assembly Containing:	69368
1	Accumet pH Meter	13-620-956
1	Beaker, 100mL	41951
1	pH-9 Bottle	31222
1	pH-4 Bottle	31223
1	Thermometer	31142
1	Support Rod	41831
1	Electrode Holder	19941
1	Battery	40367
1	Combination Electrode	13-620-108
1	Instruction Manual	69369

OPERATING CONTROLS

Prior to using the Model 956, become familiar with the location and function of each control and connector. All are located inside the case (see figure 1) and are described in detail below:

Readout Meter. An analog meter with a pH scale covering 0-14 pH units, and a millivolt scale covering = 700 millivolts. The scales are graduated in 0.1 pH and 10 millivolt increments, respectively, and are mirrored to minimize parallax errors.

Mechanical Meter Zero. A screwdriver adjustment that mechanically zeros the meter pointer. With the instrument sitting on a flat, level surface, the adjustment must be made with the FUNCTION selector in the OFF position to avoid electrical deflection of the pointer.

FUNCTION Selector. A three-position rotary switch used to disconnect battery power (OFF position) when measurements are not being taken, and to select the operating mode; pH for pH determinations and MV for millivolt measurements.

TEMPERATURE Control. A potentiometer that functions only in the SH mode to compensate for the effects of solution temperature on the electrode. The control is continuously variable from 0° to 100°C with a scale graduated in 2° increments.

STANDARDIZE Control. A multi-turn potentiometer that is used in pH determinations to set the meter to the pH value of a buffer solution, thereby compensating for the differences in the zero potential of electrode systems. In millivolt and redox measurements, the control establishes the millivolt zero reference point on the meter.

INPUT Jack. A jack that receives the BNC lead from the combination electrode, or any lead from an indicator electrode with a BNC connector.

NOTE: If an indicator electrode with a pin jack is to be used, the pin jack adapter. Cat. No. 13-620-490 should be connected to the meter.

REF Jack. A jack that receives any single-lead from a reference electrode with a pin connector.

Support Mounting Socket. A threaded socket (immediately to right of TEMPERATURE control) which accepts the electrode support rod when instrument is in operation.

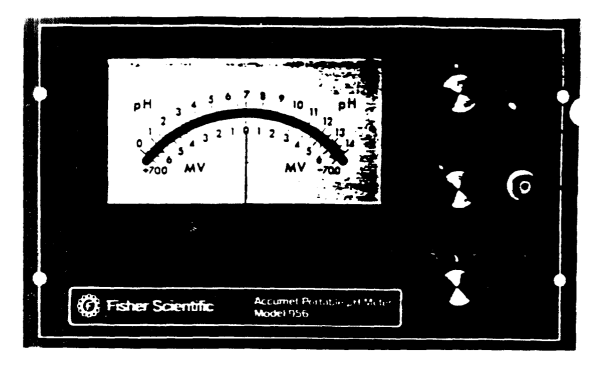


Figure 1. Controls

187335

BA TERY REPLACEMENT

The expectancy of the supplied alkaline power cell (Everady No. 522) is approximately 100 hours of total peration, if instrument usage does not exceed 4 hours/day. Low battery power is indicated by a loss of accuracy, erratic indications or no meter deflection when the FUNCTION selector is set to the pH or MV positions. A more accurate check can be made using a battery checker. When so doing, a reading of 50% (4.5) or less on a 9V battery) indicates that the battery should be replaced.

Idea. The battery should be replaced with an Eveready No. 522 alkaline power cell such as that supplied with the instrument. However, in an emergency, a standard 9-volt transistor radio battery (No. 216 or 20 may the used — but with a considerably shorter life expectancy. Additionally, if a transistor radio battery is allowed to remain in the instrument beyond its useful life, battery leakage could cause damage to sensitive metel circuitry.

Where replacement is required or when it is desired to check the battery with a checker, perform the following:

- 1. Set FUNCTION selector to OFF.
- 2. L plug electrode leads.
- 3. Firemove electrode support rod.
- 4. Firmove the four screws (2 along each side) securing front control panel to carrying case, then lift panel up and out of case.
- 5.1 snap connector, then remove battery from holding clips.
- 6. Cireck or replace battery, then reassemble instrument by generally reversing steps 1 through 5.

PRELIMINARY PROCEDURES

Prior ω making a pH or millivolt measurement, the mechanical zero of the meter must be checked and the electrode system must be properly installed.

Checking and Adjusting Meter Zero

Before initial use, the mechanical zero of the meter movement must be checked and, if necessary, adjusted. The meter pointer should indicate precisely 7 pH (or zero millivorts) with the FUNCTION selector set to OFF. If the above conditions are not met, adjust the MECHANICAL ZERO, using a small screworiver, until specified indication is obtained.

Installing pH Electrode System

The Model 956 is supplied with a plastic-body combination electrode that is especially suited to the rugged conditions associated with field work. The INPUT and REF jacks will accommodate all other indicator and reference electrodes that employ BNC and pin cable connectors, respectively. The supplied electrode holder contains openings for one or two standard size electrode caps.

NOTE: At this point, the electrode(s) should be properly conditioned according to the manufacturers instructions.

To install the supplied combination electrode, perform the following:

- Unsnap the support rod from its molded case compartment and screw (clockwise) the threaded end into the socket in the meter ponel.
- Install electrode holder (see Figure 2) on support rod by depressing plate on underside of holder, sliding holder onto support rod, and releasing plate.
- 1. Remove the plastic cot from the tip of the electrode.
- 4. Slip electrode body into slot on either side of holder and lower it until cap is firmly seated.
- 5. Observe that FUNCTION selector is set to OFF, then remove shorting cap from the BNC jack.
- 6. Insert the round BNC connector jack onto the INPUT jack.
- Check electrode system efficiency by standardizing in a buffer of known value and measuring in a second buffer by generally following the Standardizing Instrument and Performing pH Determinations procedures in the OPERATION section.
- a. The electrode is now installed and ready for use (see OPERATION section). When not in use store the electrode as directed by the following procedure.

OPL JATION

Separate procedures are provided below for standardizing the instrument, for performing pH measurements, and for performing millivott measurements. Note that in all the procedures that follow, the FUNCTION selector should always be in the OFF position when immersing or removing the electrode system from solution. This practice avoids meter fluctuations that may damage the movement. Also the FUNCTION selector must always be set to OFF when the instrument is not in use to avoid unnecessary battery drain.

Standardizing Instrument

The Model 956 is calibrated at the factory to indicate the exact pH of solutions measured at 25°C, using a signal representing the theoretical potential of a 100% efficient electrode system. However, the actual presented of each electrode system varies slightly from the theoretical and shifts the calibration point. For "reason, the instrument must be standardized prior to performing one or a series of pH determinations. The Model 956 is standardized as follows:

- 1. Set up electrode system as described under installing pH Electrode System.
- Place a buffer container directly under the electrode system, then lower electrode system into buffer solution.

NOTE: Always situate buffer and sample containers in case compartment provided to avoid damage from spillage.

- 3. Immerse the supplied thermometer into the buffer solution.
- Allow at least 2 minutes for the electrode and buffer systems to reach thermal equilibrium, then set the TEMPERATURE control to match the thermometer indication.
- 5. Turn the FUNCTION selector to pH.
- Adjust the STANDARDIZE control until the meter indicates the exact pH of the buffer as determined from a buffer pH-versus-temperature table (usually supplied by the buffer manufacturer).

NOTE: Due to the high impedance characteristics of the glass membrane, the connecting cable of the electrode should not be moved or touched while measurements are being performed; otherwise, unsuable meter indications may result.

- 7. Turn the FUNCTION selector to OFF.
- 8. Remove electrode system and thermometer from the buffer, and nose them with distilled water.

Performing pH Determinations

Proper use and care of electrodes are fundamental to achieving reliable pH determinations. Therefore, a instructions supplied by the electrode manufacturer should be carefully followed.

Assuming that the instrument has been properly standardized, a pH determination is made as follows:

- 1. Be certain that FUNCTION selector is set to OFF.
- Place sample container in case compartment directly be eath electrode system, then lower electrode system into sample solution.

NOTE: Always situate sample containers in compartment provided to avoid damage from spillage.

- Place the supplied thermometer into the sample, and allow at least 2 minutes for electrode system and sample solution to reach thermal equilibrium.
- 4. Set the TEMPERATURE control to match thermometer indication.
- 5. Turn the FUNCTION selector to pH, and read the pH value of the sample directly from the meter.
- 6. Turn the FUNCTION selector to OFF
- 7. Remove electrode system and thermometer from the sample, and rinse them with distilled water.
- 8. Repeat steps 2 through 7 for remaining samples.
- 9. Upon completing final determination each day, be certain that FUNCTION selector is in OFF position.

187383

Performing Millivoit Determinations

The Model 956 is a convenient tool for measuring electrode potential and indicating results over a =700 to +700 millivoit range. There is no need to standardize the instrument with a buffer or compensate for solution temperature. Readings can be taken as soon as the millivoit zero reference point is established, with no further standardization necessary.

Assuming that the electrodes are conditioned and installed according to instructions from manufacturer, a millivoit measurement is made as follows:

- 1. Be certain that FUNCTION selector is set to OFF.
- 2. Connect the shorting cap to the INPUT jack.
- Turn FUNCTION selector to MV position and adjust STANDARDIZE control until meter indicates zero millivolts (or any other desired value).
- Turn FUNCTION selector to OFF, remove jumper, and connect electrode leads to INPUT and REF
 lacks.
- Place sample container in case compartment directly beneath electrode system, then lower electrode system into sample solution.
 - NOTE: Always locate sample container in compartment provided to avoid damage from spillage
- 6. Turn FUNCTION selector to MV and read electrical potential of solution directly from meter.
- 7. Turn FUNCTION selector to OFF.
- 8. Remove electrodes from sample and rinse them with distilled water.
- 9. Repeat steps 7 through 10 for remaining samples.
- 10. Upon completing final determination each day, be certain that FUNCTION selector is in OFF position

SERVICE

Solid-state components and easily accessible assemblies simplify servicing of the Model 956. The following information is given to aid qualified service personnel and help minimize downtime. The information includes a brief circuit description, a troubleshooting guide, procedures to check instrument performance, buffer accuracy, electrode system efficiency, a procedure to remove the meter panel, and calibration and checking procedures. Also included at the end of the manual are replacement parts and a schematic. In addition, Fisher Scientific Company maintains a company wide instrument Service Division with fully staffed service centers to assist with any service problems.

Understanding the Circuit

A single, operational amplifier in the Model 956 circuitry provides the necessary high-impedance input as well as the amplification required to drive the analog meter. This and the use of three diodes in the reference circuit facilitates meter indication of either polarity as well as excellent battery voltage stability over the entire instrument range.

187389

If the instrument is not operating properly, select the SYMPTOM category (below) that best describes the operating malfunction, and proceed to check the PROBABLE CAUSE areas given under the category taking proper CORRECTION action as directed.

NOTE: The Model 956 is a precision, electronic instrument and should only be serviced by compa trained personnel.

Sympton	m
---------	---

- 1. Meter pointer does not respond to any **FUNCTION** switch position.
- 2. Cannot adjust meter to indicate pH of buffer for standardization.

3. Meter pointer drifts.

5. Results incorrect.

solution.

same off in any

Probable Cause

- 1. Battery defective.
- 2. Open circuit in meter.
- 3. Detective FUNCTION switch.
- 4. Defective STANDARDIZE control.
- 5. Defective On Amp.
- Electrode
- 2. Buffer defective.
- 3. Defective input circuit.
- 4. Electrode not properly immersed in solution.
- 5. No solution in reference cavity of electrode.
- 6. Amplifier defective.
- 1. Electrode defective.
- 2. Defective input circuit.
- 3. Defective FUNCTION selector.
- 4. Solution being measured is unbuffered and exposed to air.
- 4. Meter indicates the 1. pH-sensitive section of electrode cracked.
- 1. Electrode defective.
- 2. Buffer defective.
- 3. Amplifier not operating properly.
- 4. Meter defective.
- 5. Electrostatic charge on meter glass face.
- 6. TEMPERATURE Control knob misaligned.
- 7. Instrument out of calibration.

Corrective Action-

- 1. Replace battery.
- 2. Replace meter.
- 3. Replace FUNCTION: switch
- 4. Replace STANDARDIZE control
- S. Replace On Amp.
- 1. Replace electrode.
- 2. Replace buffer.
- 3. Perform all steps under Check ing Instrument Performence. if test fails, check circuit continuity with low voltage ohimmeter.
- 4. Make sure tip of electrode immersed at least one inch.
- 5. Add fill solution to reference car . ity (nongel-filled only).
- 5. Replace Op Amp.
- 1. Replace electrode.
- 2. (Same as 2 and 3 above).
- 3. Replace selector switch.
- 4. Use proper technique for me ing unbuffered solutions.
- 1. Replace electrode.
- Replace electrode.
- 2. Replace buffer. Also see Checking Buffer Accuracy.
- 3. Replace Op Amp.
- 4. Replace meter.
- 5. Apply anti-static solution to meter lace
- 6. Realign knob as directed unde Calibrating pH Scale and Temperature Control.
- 7. Recalibrate as directed under Calibration Procedures.

Checking Instrument Performance

While performing the following procedure, there should be little, if any, drift and fluctuation in the meter needle in either the pH or mV mode. The presence of drift and/or fluctuation indicates an instrument defect instrument performance is checked as follows:

- 1. Connect shorting cap to the INPUT jack. (This shorts instrument input.)
- 2. Set FUNCTION Selector to pH position.
- 3. Set TEMPERATURE Control to 25°C.
- 4. Adjust STANDARDIZE Control until meter indicates 7.0 pH.
- Turn STANDARDIZE Control to full clockwise and full counterclockwise positions; meter should indicate at least 4.0 and 10.0 pH respectively.

NOTE: This demonstrates that STANDARDIZE Control has sufficient range to compensate for the zero potential of electrode systems.

- 6. Set FUNCTION Selector to MV.
- 7. It should be possible to adjust STANDARDIZE Control until meter indicates 0.0 millivolts (center scale).

Checking Buffer Accuracy

The most practical way to check buffer accuracy is to substitute a questionable bottle of buffer with a fresh, unused bottle. Some buffers deteriorate with age, but contamination is the principal cause of inaccuracy. A common practice is to regularly use two buffers — one as a working buffer and one as a check buffer.

Checking Electrode System Efficiency

Once instrument performance and buffer accuracy are found to be satisfactory, check efficiency of electrode system as follows:

- 1. Install combination electrode
- 2. Obtain two buffer solutions that are several pH units apart in the 3 to 10 pH range.
- Standardize the instrument with one of the buffers (as directed by Standardizing Instrument procedure given in the OPERATION section).
- Determine the pH value of the other buffer (as directed by Performing pH Determinations procedure in the OPERATION section).

NOTE: The result of this determination should be within 0.1 pH units of the known buffer value. If it is not, repair or replace the electrode system.

Removing Meter Panel

Most service procedures, such as battery and component replacement, require that the meter panel be removed from the case. This can be accomplished quickly and easily as follows:

- 1. Set FUNCTION Selector to OFF.
- 2. Disconnect electrode system from meter.
- 3. Remove electrode and electrode holder from support rod, then unscrew rod from meter panel.
- Remove the four screws (two along each side) securing meter panel to carrying case, then lift panel up and out of case.
- 5. All components are mounted to rear of panel and are now easily accessible for service.

Calibration

The Model 956 is factory calibrated to perform within the stated pH and MV specifications. Under normal conditions, this calibration will yield satisfactory results over a long time period. However, it is recommended that the instrument be recalibrated annually or after any circuit component (except battery) has been replaced. An internal component view of the meter is shown in figure 2.

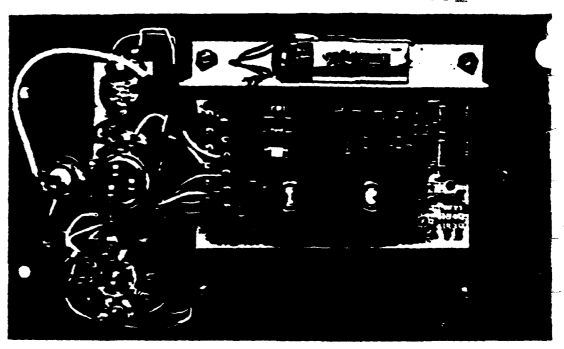


Figure 2. Internal Components

Calibrating Millivolt Scale

The millivoit scale is calibrated as follows:

- 1. Perform steps 1 through 3 under Removing Meter Panel.
- Remove the four screws (two along each side) securing meter panel to case.
 NOTE: For calibration purposes, it is not necessary to remove panel from case since calibrating putometers are accessible by lifting panel up from left side.
- Check and, if necessary, adjust the meter mechanical zero as directed under Checking and Adjusting Meter Zero.
- 4. Connect the supplied shorting cap to the INPUT jack.
- 5. Turn FUNCTION Selector to MV position.
- Adjust STANDARDIZE Control until meter indicates exactly 0.0 mV, then return FUNCTION Selector OFF.
- Remove cap from INPUT jack and in its place, connect a millivolt source (with accuracy of 0.1% or hetter)
- 8. Adjust millivolt source to produce an output of +500 mV.
- Model 956 should indicate + 500 mV. If not, readjust R2 on circuit board until + 500 millivolt indicato
 is obtained.

NOTE: Positive deflection of meter is to the left of the 0.0 center position.

- 10. Reverse polarity of millivoit source.
- 11. Model 956 should indicate 500 mV. If not, readjust R2 until the meter indicates midway between 500 mV and the incorrect negative reading.

Calibrating pH Scale and Temperature Control

The pH value of a solution is generally dependent upon temperature; hence, the accuracy of a pH scale is dependent upon the accuracy of the TEMPERATURE Control. Consequently, both these components are calibrated in conjunction with each other as follows:

NOTE: It is recommended that the mV calibration procedure be completed before calibrating the pH scale. Therefore, the procedure given below assumes that meter panel is not secured to the case and that the millivoit source is still connected to the INPUT jack.

- 1. Set millivoit source to 00.00 or disconnect source and connect shorting cap to the INPUT jack.
- 2. Turn FUNCTION Selector to pH and, if necessary, readjust STANDARDIZE Control until meter indicates exactly 7.00 pH.
- Rotate TEMPERATURE Control to full clockwise and full counterclockwise positions. Knob pointer should approximately align with the 0 to 100°C scale markings. If not, loosen the two knob setscrews and realign knob until above specifications are met.
- 4. Set TEMPERATURE Control to 25°C.
- Connect millivolt source to INPUT jack (if disconnected in step 1) and adjust millivolt source to supply an output of + 295.785 millivolts.
- 6. The meter should indicate 2.0 ±0.3 pH. If not, adjust R10 on circuit board until correct indication is obtained.
- 7. Reverse polarity of millivoit source.
- 8. Meter should indicate 12.0 ± 0.3 pH. If not, readjust R10 until meter indicates midway between noted reading and correct reading.
 - NOTE: If desired, Model 956 pH scale can be calibrated using buffers. In this case, instrument should be first standardized with a pH 7.00 buffer and then calibrated using a pH 4.00 and adjusting R10.
- 9. Reassemble instrument by generally reversing steps 1 through 3 under Removing Meter Panel.

Replacement Parts

The replacement parts, along with their location drawings, are provided in this section. The section is arranged in assembly to subassembly breakdown. That is, the highest order assembly appears first, followed by its subassembly make up. Non-field repairable is semblies or subassemblies are not broken down into their component parts. Note that parts information is only valid at the publication date (see back cover) of this manual, and subsequent revisions may have occurred after publication.

TAB

APPENDIX E

YSI MODELS 33 AND 33M S-C-T METER INSTRUCTION



MAL MACRITURE

The TSI Model 33 and 33E 3-C-? Meters are portable. bettery povered, transporterized pastruments designed to securately measure salinity, readestivity and temperature. They use a probe emalating of a respect plantic conductivity call and a processon TEE thornsons tumperature sensor mushand to a single wait.

Conductivity with the Endel 33 is empressed as emerg-aboutcontinuous: with the 33%, it is millisteness! meter (mi/h). These are measurements of the electrical contactance the sample would show if measured between eppeats face of a 1 on cube. (Conversion informations 1 sicrembe/on = 0.1 of/m.) Salinity is the number of grans of sali/kilogram of sample (ppt-= parts per thrushed). This measurement assumes the sample con-tains a "standard" one veter sult mixture. The sample temperature is measured in degrees Calsius.

Salisity measurements are memally temperature compas-sated by direct dial. Community measurements are not temperature companated; however, a temperature function is provided on the instrument to aid with extendation of excreptions. Then temperature and exceptivity are known, it is possible to extend to sale. inity; and whom only temperature and calimity are known, it is possible to calculate conductivity. This is discussed in the section on localibration.

CHETTERTIES

Brief 33 Conductivity

Easess: 0 to 900 (EL), 0 to 5.000 (EL0), and 0 to 54.000 micrombos/on (miss) with TSI 3300 Series Frebes. (Bote: The "mistered" designations on the meter are a shorthand form for "microshe/en".)

Accuracy: (See Error Section) \$2.5% max, error at 500, 5,000 and 50,000 plus probo-51.00 max, error at 250, 2,500 and 25,000 plus probo-

tondability:

2.5 aucressos/es es 500 aucresso/es rases. 25 ELECTROMOS/OR OR 5.000 ELECTROMO/OR CAMPIL. 250 ELECTROMOS/OR OR 50.000 ELECTROMO/OR CAMPIL.

Tonnerstore Consessation: Rese.

Bodel JEE Condustivity

0 to 50 (ml), 0 to 500 (ml0), and 0 to 5,000 in ma/a with TEI 1300 Series Probes.

LOURSHY: (See Error Section) \$1.30 max. orrer at 50. 300, and 5,000 plus probe. \$1.00 max. orrer at 25, 250, and 2,300 plus probe.

Loodability

1.25 mi/s on 10 mi/s rance. 1.5 mi/a es 100 mi/a ranço. 25.8 mi/a on 5.000 mi/a cango.

Temperature Commencation: Sees.

Salisier

Ranger 0-40 ppt in temperature range of -2 to -45 Within specified conductivity range of 0 to 50. Seculibration.

Assuranty (See Error Section)
Above 4°C: g0.9 ppt at 40 ppt and g4.7 ppt at 20 195 1km en MEETLYSTY proce.

balow ("C: 21.1 ppc at 40 ppc and 20.5 ppc at 10 pot plas conductivity probe.

handahility: 0.2 ppt us 0-40 ppt reaso.

Pemperature Compensation: Essent by direct dial =2 to +45 C.

Temperature Range: -3 to +40°C.

hommory: 40.100 at -200, 40.600 at 4500 plus : (See Street Section)

Bendahility: 40.15°C at -3°C to 40.37°C at 45°C.

Power Supply: Two D-cise alkaline betternes. Fre 295 or equivalent, provide approximately 200 bec CONTRACTOR.

Instrument lablest lange: -1 to $+15^4 C$. A same of 90.15 of the reading per $^4 C$ charge in last: erature can occur. This error is socilotable ! instrument is reedjusted to redline for each read



YSI Scientific

ow Springs, Ohio 49367 USA - Phone 913 767-7241 - 800 343-HELP

THE 1300 Derive Conductivity/Temperature French

Project Frebe Constant: E = 5/cm (E = 500/a)

ASSESSED : 270 of reeding for conductivity and salinity.

Breef of 14.100 at 600 and 14.100 at 400c.

CHARTE

letm

- 1. Migust meter zero (if somesary) by turning the hability seror on the meter face so that the meter social emissions with the zero on the conductivity scale.
- 2. Calibrate the secor by turning the MBE control to REDITE and educating the REDITE control so the actor seedle lines to with the redites on the actor face. If this cannot be accomplished, replace the batteries,
- 1. Fing the probe late the probe just on the side of the instrument.
- 4. Pet the grobe is the solution to be measured. (See Probe Wes.)

?mpereture

Set the SUE control to TENTERITUE. Allow time for the probe temperature to some to equilibrium with that of the water before reading. Boad the temperature on the bettem scale of the meter in degrees Coleius.

Conductivati

1. Switch to Eids. If the resting is balow to on the 0-100 range (5.0 on the 0-10 ad/a range), switch to 210. If the resting is still below 50 (5.0 ad/a), switch to the Eidself. Send the nature scale and maintainy the resting appropriately. The assure is oppressed in microscotom (as/a). Headurements are not temperature compensation.

ماحصط

Botor Souding: 347 (34.7 mg/m)

Seale: Il9

lasver: 2470 misreshee/en (247.0 mi/n)

2. Then asserting on the MISS and MIS senior, toproce the CML THIT betten. The motor recting asself fall loss than Mr if greater, the proce is isolat and the seasonness is in error. Clean the proce and reseasons.

MOTE: The CRL TEST does not traction on the 21 scale. MINIST

Salinity

- 1. Seturnize the sample temperature and adjust the temperature dial to that value.
- 2. Switch to gion. If the reading is above 100 microshe/on (50 ms/m), the salinity value is beyond the measurement range.
- 3. If the reading is in range, switch to SALIKITY and read salisity on the red 0-40 ppt meter scale.
- 6. Deprese the CEL TEST button. The fall in meter reading should be less than 20: if it is greater, the probe is found and the measurement is in error. Clean the probe and re-measure.

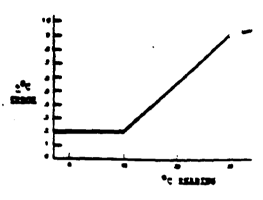
Bre

The maximum error is a reading can in the fellowing section.

Tenserature from

The temperature state to designed to the salitate experience restates compensate salitates measurements.

Pieure 1 shows total error far proce as verses of motor reading.



F16012 1

Masple: Mar Reeding:

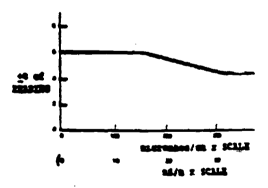
15°C

Total Server

13°C 90.4°C for prote /

Conductivity Serve

Figure 2 shows the worst-case enducts to tunction of the conductivity reading it is instrument continued.



Emmple Enter Seeding: Seale: & Anoting Error: Longrapy:

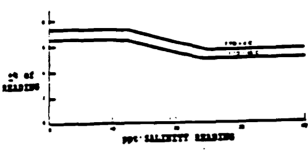
360 misroshoo/en (36 mis-230 24.50 3600 e162 misrosh /c mi/m) tor probe ab* 35'

77000E 2

TALLASTY BYEN

The solicity realizes are a function of temperature and eccelerators, therefore the assurery is a function of both.

The temperature settle and temperature control here been designed to minimize the temperature error contribution to the califacty error. The error shown in figure) is the cotal of the temperature and conductivity probe, the temperature peaks and the salinity scale error.



77.00E 3

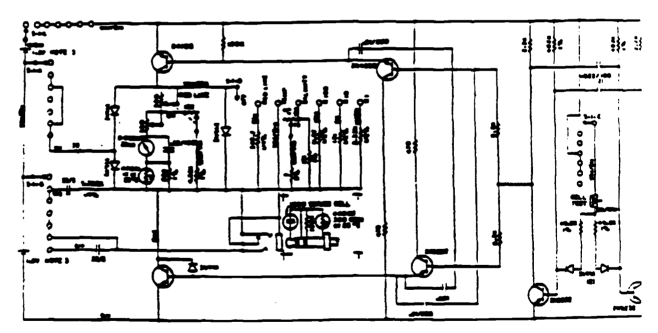
Example
Refer Resilies: 10 ppc. 0 10°C
% of Resilies Error: 6.30
Lecurory: 10 ppc -0.65 ppc for all errors.
Lecurory: case.

The circuit is composed of two parts: a emittripreter and systemate translators. The emittripreter produces a square veretors voltage. The openry were is applied to two systemate translators. They alternately applied to two systemate translators. They alternately applied to two systemate translators. They alternately applied two battery as providing at power value eminists polarization of the centre of th

DESTRUCT MATERIALS

The only maintenance required is battery replacement. Two "F" size alkaline flashlight cells, such as Evereasy EDS or equivalent, will provide 300 hrs. of operation. Accuracy will not be maintained if time-restrict
"F" cells are used. Bettery replacement is indicated
when the redline adjustment cannot be occupilized.

Seplace batteries every six mentls to reduce the descriof corrector due to looky batteries. To replace satteries, remove the serves from the rear cover. The battery balders are color coded. The positive end sust go on red.



SOTES: Resistance values in chas. I = 1,000. tesisters are 1/2 F. 100. unless etherwise specified.

Sattery is 9 size. alkaline only. Everency 8-95 or ornivalent.

This schematic is representative and may be slightly different from the elecute in year instrument.

THE MOSEL IS AND USE SCHEMATE

read the salinity for this sample.

Recalibration should be done at the factory. If concessory contlibration is accessory, however, one of the precedures described below may be attempted.

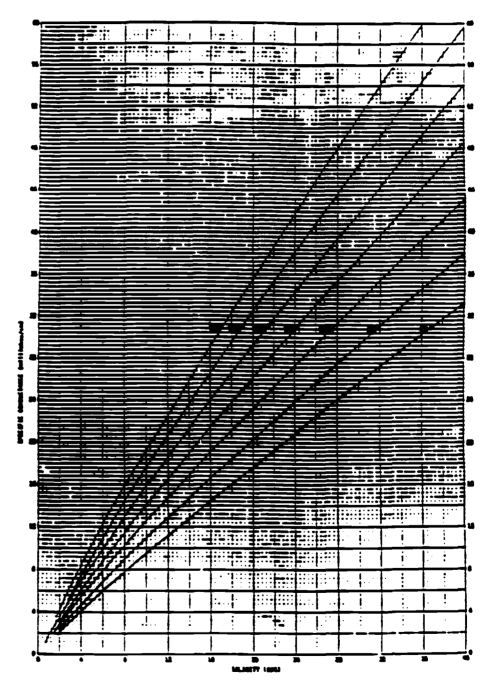
1. One this author if the temperature than become lesses or suppost from its sornal position.

a, fined the temperature and conductivity of 4 sample. Determine the salinity of the selection by running a line vertically on the Calibration Graph from this conductance value until it intersects the appropriate of line (interpolate as required for temperature between the given of lines). From this intersection animal a line berimmtally to the edge of the graph to

Stample: 25.000 aterestes/en (2.500 at 1: 51700 a salinate of 17 ppt.

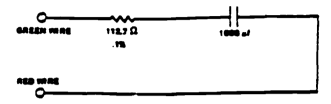
b. Remove the temperature tash, grates to 27 turn the course shaft until the geter seedle . the salinity value determined in Step 4. In amoin given, the value is 17.

c. Switch to TEMPERITUE and note the reading must be the same as Step in a: 1/ it began again. Replace the mass switchest ur control shaft) with the pointer at the same persture as the noter reading and tighten : screwe securely.



- 1. Too may use the recister and expected health shows in the shores to constitute for the process in the following recalifration processes.
- a. Set the instrument for a salinity seasurement as sermal.
- b. Substitute a 1000 exercisrat capacitor and 112.7 one 0.10 telerance recistor for the proce.

Consert the reclater and espectter between the green vire and red vire on the just consections inside the instrument.



- c. Turn the temperature dial until the meter reads rediine.
- 4. Reinstall the temperature tack with the arrow at $15^{\circ}\mathrm{C}_{\star}$

This is a temporary exhibitation only. Seturn the instrument to the factory for proper recalibration.

TER 1300 SERVER COMMICTATION/COMPERATORS PRODUCT

Becartettes

These probes are designed and constructed for requisions probes as field use. The conductivity $\epsilon_{\rm c},i$ constant is 5.0/em (§60.0/m) giv. Each probe contains a processes TSI thermister temperature sensor of $\pm 0.1^{\circ}{\rm C}$ accuracy at 0°C and $\pm 0.1^{\circ}{\rm C}$ at 40°C. The low capactions cable assembly terminates in a three terminal 0.23° dis. phone plus.

The 1310 has a 10 feet cable and the 1311 a 90 feet cable. Other lengths are available on special order.

The probe has a rigid P.Y.C. body, platinized pure bielel electroles. and a respect cable.

Cleaning and Storage

Classias

Then the cell test indicates low readings the probable cause is dirty electrodes. Eard veter deposits, cals and organic natter are the most likely contaminants.

For convenient normal cleaning seek the electrodes for 5 minutes with a locally symilable betaroon tile cleaning properation such as low Chemical "lathroom and Chrone Cleaner." Johnson Fax "Envy. Instant Cleaner." or level Brand "Basia. Tab. Tile Cleaner."

For stronger cleaning a 5 minute souk in a solution made of 10 parts distilled water, 10 parts isopropyl sichel bid 1 part SCI can be used.

livery riase the probe thereoghly in tay vater, then in distilled or described vater after cleaning and before storage.

CLUTTON: Do not touch the electrodes inside the probe. Platinum black is soft and can be scraped off. If cleaning took not restore the prese perform

Sterage

It is best to store conductivity coils in delet veter. Calls stored in veter require less from platinization. May coll that has been stored should be sented in descriped veter for 14 hours to use.

Real Attackment

- 1. Close the probe.
- 2. Place the cell in a 10 ml (approximate) jabeater and add ecough TEL 1140 Platinizing Solution cover the electrosis. Do not cover the top of grade.
- J. Fing the probe into the Godel 33 or 13E. swit the E100 scale to platinize the electroses. Here probe slightly to obtain the highest meter reading continue platinizing for the approximate time is

Boter Booking	
15/1	fine i.
1.000	5
2.500	i
1.000	ì
1.500	11
1.000	16
	35/8 3.000 2.500 3.000 1.500

- 4. After the elapsed time, remove the press and r in two exter, then in distilled or desembled vater.
- 5. Seturn the solution to its container. 2 or solution should be sufficient for 50 treatments.

Proto the and Promotions

- 1. Obstructions near the probe can disturb read at least two lastes of chearance must be allowed non-motallic underwater objects. Betailic objects as place or wrights abould be bept at least least the grants.
- 2. Verghts are attached to the cubic of the TII and JILL Probes. The TII JI27 Verghts are explice pairs with a total verght of 4 camers per pair. In it become secretary to add note verght to every veter currents, we support limiting the total we to two passeds (8 pairs). For verghts in agrees of passeds was an independent purposesses cable. In excess, verghts must be bopt at least 6 taches every the probe.
- 1. Sentia agitation by raising and lovering the poweral times during a measurement insures flor specimen solution through the probe and improves time response of the temperature sensor.

Conductivity and falinity Corrections for Long Cabl

The editional leafth of wire in leng cables expectance and registance which will effort read: The recommended very to correct for tease influence by use of TEE Conductivity Calibrator Solutions below!, which will permit an estimate of correct factors. If these solutions are not evaluable. Soliowing tables can be used for the correction errors council by cable resistance and capacitanc operal length versions of the 1916. 1911. 5-1791) 3-16120 probse.

187400

Cable Leagth is feet Indicated Conductivity 200 300 50 100 500 1000 Lance who/es 15 -1.0 -1.0 重 -2.0 -1.0 -4.8 4.0 100 4.1 -1.2 4.0 100 21.00 -0.6 -1.0 4.0 4.1 . 4.1 4.5 -2.0 -3.0 -1.0 فله -3.0 4.0 1000 4.66 4.3 4.6 -1.2 -1.8 4166 4.01 4.05 4.1 4.2 4.3 4.5 -1.6 21.00

" This row indicates the effect of the chance from 100 Es to 600 fr wom the instrument is evitable to the 210

IR represents conductivity levels thick to do not recommend for produce with the indicated cable lengths.

TABLE 17: THEFENTINE COMMETTICES IN C.

Indicated Temperature		Cable	Langt	h in P	eet .		
46	18	50	100	200	300	100	1000
0 18 20 40 50	FOR	2016 1016 2016 2016 40.3	40.1 40.1 40.3	40.1 40.2 40.3 40.4	4.3 4.4 4.7	40.3 40.5 40.7 41.1	•0.6 •1.6 •1.4 •2.3

TABLE ITT: SALIETYT COMMETTICES IN PARTS PER TROUBLES

Temporature Setting	Continues of the contin	Cörroetians	tres table !
•€	-1	~4	-18
ŧ	-9.1	-4.5	-1.0
36	1000	4.1	→.1
20		-4. 3	4.6
9	3000	-1.25	→.5
		→.2	4.4
50		4.13	-4.25

BOTES:

- 1. CHARACTERICO CONTROPORTIONS Should be made true Pable I. or by exterioletion of the table.
- Isliaity environtiese regules determination of enedescripts, bear conjustivity correction, and the set-ting of exercised temperature readings prior to sal-LILEY SOLDERSBEERS.
- i. One of these commercians should increase the error bers tos emperentes by less then 100.
- 4. If your expressive conditions are such that a 20 se greater employety of correction to required, the soll toss feature vill not properly indicate a defective

Call Calibration and Standard Solutions

The seal constant of a conductivity call day rest slightly such the conductivity of the colution being consumed. Soll calibration may also be affected by

exections imming, regularization of or spect. A coul and never can be callers. I t a system. Them TSI 3160-3169 Commetitity Selutions.

THE Conductivity Calibrator Sciences ** " a full tempered discussion and detr for see.

Part Dunber	\$1.00	Conductivity at 2 .00
TEI 3161	Galles Gaart	1000 atereme/es -4.5
TRE 1163	falles Graff	18.000 atereme/4 # 18.000 atereme/4 #
TEI 1164 TEI 1165	falles feart	100.000 sterento/en :
TEI 1167 TEI 1168 TEI 1169	l Piats l Piats l Piats	1000 atermite/es 10,000 atermite/es 21

Directions for calibration at temperatus a 18°C are included with the Canadactive p talestere.

In calculating the cell constant in terms, the uncertainty of the motor ca br ter feleties.

THE MINE, 13 AM 134 WAS THE THE SIA. "1.

STATE indicates that the corrections are less than \$.10°C. If the salinity measurement is to be used for correction on the 512. the reading should be to Calernetty. The termils is:

PPE Calmentar . ((Salinity ppc -0.8) 11...8)

for those instruments the 0.03 cm in project equation simplifies to:

PPE/Cl = (solinity in set z 18^3) /1.8

For solimity correction when using the Bortal no the salinaty resting tirest from the in JEE. No conversion is necessary.

Model 33 and 328 calinity rectings taken in c with model 54 disselved expres reclines e | a t has presides was be learn end to essentiable salinity extractions to dis it desse. Correction tables are evaliable from the

TARREST !

111 THE products carry a con-year variably of skip and party, amminute of betteries. Dead sortdoot, microse, or temperine will be 1"94 mand cherry.

If you are experiencing difficulty with any 17 it may be returned to an extheriset 731 repair, even if the varranty has espired. If tersory engletance for any reason, contact

Product Service Separtment Teller Springs Instrument Co., Inc. 1725 STREETS LANG 7.0. bon 379 Tellow Springs, Ohio 45387, 8.3.4. Phone: (513) 767-7241 (600) 343-8829

YSI Scientific

"Mary Springs, Chap 45567 LEA + Phone 515 767-7241 + 600 545-466.5

Part Dusber 103309 : Item 021470

TAB

APPENDIX F

187403

. TABLE OF CONTENTS

	_ FARS
1.	Specifications
II.	List of Standard Accessories
III.	Pre-Operation Check Out2
IV.	Operation and Description
ν.	Recorder Output
vI.	Critical Heasuring Area4
VII.	Calibration Procedures
vIII.	Trouble Shooting8
ıx.	Maintenance
x.	Parts & Accessories
XI.	Warranty

DET TURBIDIMETERS

OPERATING INSTRUCTIONS

187404

POREMORD

The DRT Turbidimeters are continuous reading Nep...l instruments which measure scattered light from particles in s s in a liquid. The optical signal is stabilized and amplif energize a meter. The instruments use solid state elecomponents and resist thermal variation and lamp aging.

All modes of the DRT Turbidimeter Instruments provide display of turbidity in Nephelometric Turbidity Units. Note the (Jackson Turbidity Units), FTU (Formazin Turbidity Units) as (Nephelometric Turbidity Units) are comparable.

The material contained in this manual will help the users : full advantage of the instruments in the majority of appl of However in event that unusual circumstances or problems not o we this manual arise, please contact your local distributor of manufacturer.

HF scientific, inc. 3052 Metro Pkwy, SE Fort Myers, Florida 33916-7539 Phone: (813) 337-2116 FAX: (813) 332-7643

Our engineering staff is available to help you with your apneeds.

187405

I. SPECIFICATIONS FOR DET-15 C

2 Ranges: 0-20, 0-200 NTU Ranges NTU (+ or -) 1% of Full Scale on either range Linearity Repeatability (+ or -) 1% of Full Scale on either range Sensitivity 0.01 NTU change on "20" range Response Virtually immediate in all ranges Power Supply 6 Volt battery. 2.6 amp hours 115/230 VAC. 50/6 Battery Charger Controls Combination Range Switch for: ON/OFF Range Select Reference Adjust Recorder Output 0-1mA adjustable, 100 maximum resistance Reference Standard 0.1 NTU (Nominal) Dimensions 11" x 9 1/4" x 5 1/2" $(27 cm) \times (22.7 cm) \times (13.5 cm)$ Weight 4.5 lbs.(2.05 kilograms)

II. 'IST OF STANDARD ACCESSORIES

QUANTITY	CAT.	DESCRIPTION
1	50083	Instruction Manual
1	60002	Reference Standard 0.1 NTU (Nomina
1	70048	Battery Charger 115 V 60 Hz OR
	70020	Battery Charger 230 V 60 Hz
2	50050	Cuvettes complete with screw top
1	50009	Light Shield
1	70825	Recorder Plug

والمتعلق والعادل بعاده والماري والتناوي والتاريخ

Extreme care should be taken when handling the Reference it or sample cuvettes as surface scratches or finger saudges cause analysis errors. Handle these items by the top on ...

Turn the Range Switch on to either range position in disconserve the condition of the battery. When sufficiently of the red low battery light will be OFF. If the light is battery should be charged a minimum of 12 hours before using instrument on battery power.

The battery, when new, usually requires several cycles f charging and recharging in order to obtain optimum rated between charges. NCTE: The turbidimeter is not meant to c; from the charger. If the battery does not have a su fi charge, the red low battery light will be OFF.

The turbidimeter provides up to 4 1/2 hours of continuous c tion as a portable battery operated unit between recharge.

This unit does <u>not</u> require any warm up time before taking ings. It is recommended that the unit be turned off be readings in order to obtain longer battery life between to es. If used as a stationary unit, leave the charger plugged This will keep the battery at an optimum level at all times.

IV. OPERATION AND DESCRIPTION

The EPA recommends that cuvettes used for instrument callb or sample measurement be matched or indexed.

For accurate measurements in the low range rotate the cuvet the well to obtain the minimum reading. Mark the cuvette of the adhesive dots provided and the instrument so that orient of the cuvette will be identical each time it is placed instrument.

To operate the turbidimeter, switch to the "20" range and the Reference Standard (0.1 NTU) in the optical well.

With the light shield in place over the well, adjust the fence Adjust knob to cause the meter to read the reference dard value on the scale. The unit is now ready for use in angle.

To make a measurement of a sample, clean one of the cuvette: fill to within approximately 1/2" of top with the sample. the top on the cuvette and carefully clean the outside surfathe cuvette with a lint free wiper such as Kimwipes. Place sample in the well and place the light shield over the Select the appropriate range for best readability.

If the instrument has been subjected to cold (below 10 de Celsius) and then brought indoors, it should be allowed to up before use, since condensation may form on the various le This can be aided by leaving the case open and the instrument for approximately a half hour.

V. RECORDER OUTPUT

The DRT-15C is shipped complete with a 0-1 mA Recorder Ou The jack is located on right side of the chassis (refer to J figure 2). To use, connect the 1/8" miniplug provided to recorder. Adjust R11 to obtain a full scale output compatible full scale reading on the DRT-15C. Once this adjustmen made, the DRT-15C will always be set up for this recorder.

VI. CRITICAL HEASURING AREA

The critical measuring area of the sample containers is the wide band starting 5/8° above the bottom. Keep this ar and free of scratches or abrasion. Handle by the top r (See Figure 1).

GLASS SAMPLE CONTAINER 28 MM O.D.

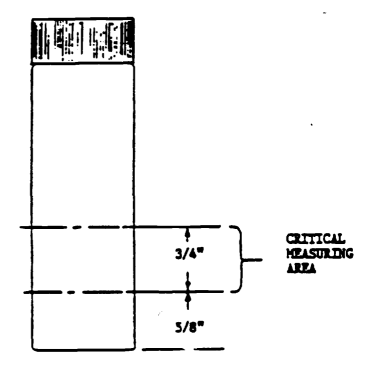


Figure 1

A. STANDARD FORMAZIN SOLUTIONS

Calibration of this instrument is based on Forma material which can be made by synthesis and reprepeatedly within one percent. When properly mixed, uniform in the number, size and shape of its particle making it an ideal turbidity standard. The unit of making it an ideal turbidity standard. The unit of making it an ideal turbidity standard. The unit of making it an ideal turbidity standard. The unit of making it an ideal turbidity standard. The unit of making its instrument is based on Formazing material which can be made by synthesis and representation of its particle making it an ideal turbidity standard. The unit of making its instrument is based on Formazing material which can be made by synthesis and representation in the property mixed.

Calibration samples may be obtained by diluting F stock suspension using "Turbidity-Free" water. F stock suspension may be prepared by the user (Re A.W.W.A. "Standard Methods", 14th Edition) or it purchased in kit form, HF scientific part number 5004

Each kit contains:

- -- 1 liter of 4000 NTC Stock Suspension
- -- 1 Gallon (3.79 liters) turbidity-free water
- -- 7 Sample cuvettes (28mm), with screw caps
- -- Instructions for dilution
- -- 1 200 µl Pipette

The following table gives the recommended dilutions stock suspension. Re sure to adequately mix the suspension prior to removing a portion for dilution.

Formazin Standard 200 ml flask and dilute \ m

Pipettes Required in NTU's with "Turbidity-free water"

9ml & 1ml in 1/100 198 9.9ml of 4000 NTU stock suspense of 400 NTU formazin dilute in 1/100 20 0.95ml of 400 NTU formazin dilute \ m

The equipment recommended for the above are: 1-1ml in 1/100 TD pipette, 1-9ml TD pipette, 3-200ml volumetric The 400 NTU formazin dilution can be made from 1 10:1 dilution a 4000 NTU stock suspension and should be made prior to making dards.

NOTE: 1. When the prepared samples start to flocculate, they are unreliable and fresh ones must be made. This will occur a rapidly for the lower value diluted suspensions.

2. The value of "Turbidity-Free" water is approximately 0 This value has been added to low value dilution, i.e., 2.1 includes 0.1 NTU for water.

SOLUTIONS 187410

The DRT Turbidimeters have been carefully calibrated be factory. However, should the Electronic P.C. Bo re photo Detectors, or the Light Source be replaced o descripted formation suspensions indicate a expectation, this may be easily accomplished in facility.

To carry out a complete calibration the following fo suspension values are required:

198 NTU , 19.8 NTU 4 2.0 NTU Fill, cap and label a separate cuvette with a sample of each.

Always mix the contents of each cuvette by inverting several times before placing in the Optical Well for a reading.

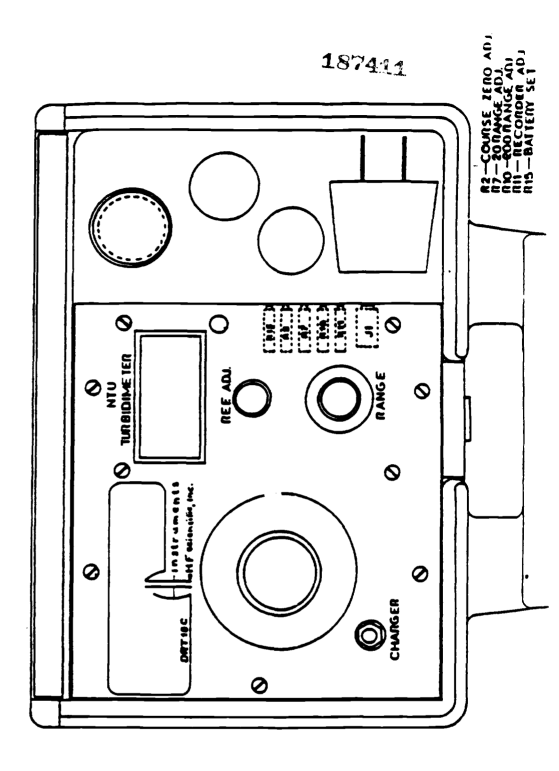
Reep the outside surface of cuvettes clean.

When placing any standards in the well, always use the Light Shield to cover the well in order to ke pout ambient light.

To gain access to the trimpots, remove the accessories from foam holder. Refer to figure 2 for trimpot identification of the new few steps.

- 1) Center the reference adjust control on top of the name.
- 2) Insert the reference standard and turn the range contro the DRT-15C to the 20 range. Adjust the "Course triapot (R2) until a reading of 0.10 NTU is obtained
- 3) Replace the reference standard with the 19.8 or standard and adjust the "20 Range Adjust" trimpot (R7 obtain a reading of 19.8 NTU ± 0.1 NTU.
- 4) Replace the 19.8 NTU formazin standard with the restandard and adjust the reference adjust control to cotreading of 0.10 NTU.
- 5) Repeat steps 3 and 4 until no further adjustmer's required.
- 6) Turn the range control on the DRT-15C to the 200 rangert the 198 NTU formazin standard and adjust the Range Adjust" trimpot to obtain a reading of 198 ± 1 "Ti

This completes the calibration of the DRT-15C.



Sympton

Heter does not respond when a sample is set into the well

Possible Cause

- 1. Lamp is burnt out. Lamp should be replaced.
- 2. Printed Circuit Board faulty. Replace Printed C: Board.
- 3. Battery is dead. Replace battery.

In the case of 1 or 2 the instrument should be recalibrated. lamp is an exceedingly long life lamp and therefore replicis infrequent.

Sympton

Reference Adjust knob does not have enough travel to adju t the reference standard value.

Possible Cause

- 1. Scratched or rubbed reference standard container or reference standard. Replace the standard.
- 2. Optics have aged. Recalibrate.
- 3. Faulty lamp. Replace the lamp and recalibrate.

Symptom

The display will not stabilize when the reference standard is the well.

Possible Cause

- Light shield is incorrectly placed over the well.
- 2. Battery has lost its charge. Low battery light is Ol battery requires recharging. When the battery is charged, the voltage will drop off causing the meter disto drift.
- 3. Cold sample causes condensation on cuvette which affec a light reading being received by the detectors.
- Unit has not been given sufficient time to stabilize ambient temperature conditions after a change of tep ture.

IX. MAINTENANCE

The DRT-15 is not designed for field servicing. It shows that the service requirements. The exceptions are Battery are Assembly replacement. This can be done in the field provinew battery is hooked up in the same manner as the Battery replaced.

A. BATTERY REPLACEMENT

Should the battery (HF catalog #70006) fail, it can be replaced. Make certain the instrument is turned off, all the accessories in the foam holder. Next, remofoam holder by placing fingers in the cuvette holpulling the rear of the foam forward, then up. The phillips head chassis screws should be be removed, the chassis from the case by sliding the chassis all to the right then pulling up. The two screws on the side of the chassis need to be removed now. This will for the removal of the battery clamp and the old b. Replace the battery by reversing the procedures desabove. When connecting the battery, be certain the connect the red wire to the terminal marked (+) as black wire to the terminal marked (-).

B. LAMP REPLACEMENT

The lamp source (HF catalog \$21084) used in the DRT-1: an extremely long life. Before replacing the lamp certain that the battery is not in need of a charge a not defective. To replace lamp remove the chass described in battery replacement. Remove the lamp will TB1 terminals numbers 9 and 10. To remove the lamp, the two set screws on the outermost barrel with a 5/6-wrench and pull the lamp straight out. Replace the latter reverse order. Make sure the lamp is pushed all thin. The set screws should be snugged up; excess precould damage the lamp. Replace the chassis in the cas recalibrate as described in calibration procedures.

I. PARTS & ACCESSORIES

HF scientific, inc. 187414
PARTS AND ACCESSORIES FOR DRT-15 C

CATALOG NO	DESCRIPTION
60002	Reference Standard 0.1 NTU
50083	Instruction Manual (DRT15C)
50051	Cuvettes - Screw Top, 3/pk
21084	Lamp Source Assembly
70006	Battery - 6 volt
70048	Battery Charger - 115 volt
50009	Light Shield
20850	Photo Diode
70020	Battery Charger, 230 Volt
20742	LED Display
50076	Printed Circuit Complete
50040	Formazin Stock Solution Kit
70900	Cuvette Cleaning/Conditioning Solution
	HF scientific, inc.
	3052 Metro Parkway, S.E.
	Fort Myers, FL 33916-7539
	Phone: (813) 337-2116
	FAX: (813) 332-7643

TAB

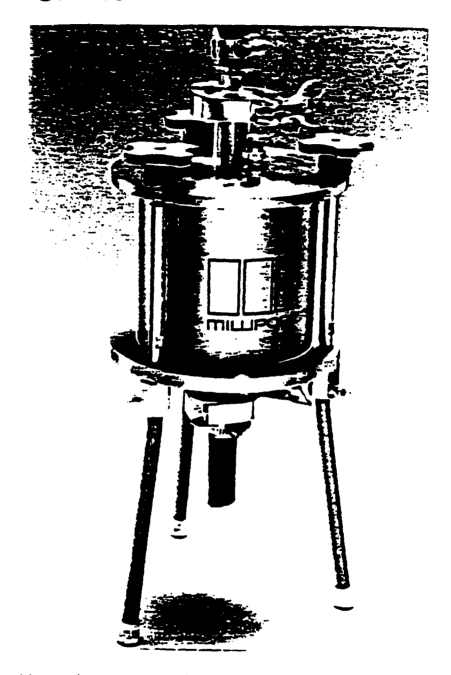
APPENDIX G

APPENDIX G

MILLIPORE

OPERATION AND MAINTENANCE INSTRUCTIONS

Operation and Maintenance Instructions 187417 OM100



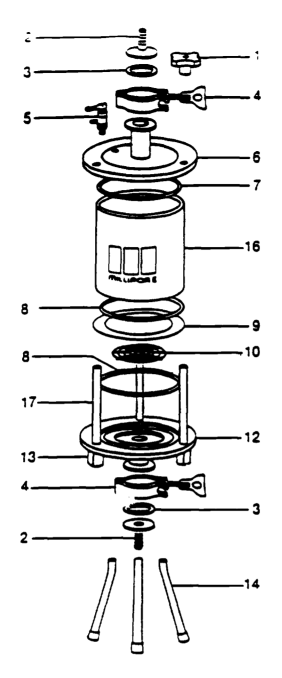
Hazardous Waste Filtration System Cat. No. YT30 142 HW



INTRODUCTION

The Millipore Hazardous Waste Filtration System (YT30142HW) is used in the toxicity determination of a solid waste in compliance with the Resource Conservation and Recovery Act (RCRA). This system has been designed to allow for absolute filtration and separation of solids from liquids by pressure filtration. The 1.5 L extension barrel; top and b. tom plates; underdrain support; filter support system; relief valve; and connectors are all construct of stainless steel. To avoid extraneous contamination from the holder assemblished, all surfaces of the components (exposed to the test sample) are Teflon coated. The TC connections of the inlet/outlet of the system allow for ease in removal, cleaning, and maintenance.

187419



	COMPONENTS	C	ATALOC
1.	Handwheel Knob (3)		YY22]
2.	T.C. Hose Connector (2)*		340
3.	T.C. Silicone Gasket (2)	10/pk	4420C
4.	Tri-Clamp [®] , 1 1/2" (2)		4Y20C
5.	Vent Valve*		331
6.	Assembly Top Plate*		
7.	Silicone O-Ring	4/pk	/Y221
8.	Teflon O-Ring (2)		YY22:
9.	Screen Support, 142mm.*		YT40:
10.	Underdrain Support*		YT301
11.	Allen Hex Key, 3/16 (not shown)		••
12.	Assembly Bottom Plate*		
13.	Leg Bushing/Socket Screw (3)	3/pk	YY221
14.	Handwheel Wrench (not shown)		• •
15.	Literature, O&M (not shown)		YT30:
16.	Cylinder, 142mm.*		114
17.	Cylinder Extension Post (3)	3/pk	XX420
	*Teflon-coated		

Hazardous Waste Filtration System Cat. No. YT30142HW

			_		TERE
AVA	11.	ARI	F	FII	TFDS

HAWP14250	0.45um MF-Membrane, 142mm (pk/50)
DAWP14250	0.65um MF-Membrane, 142mm (pk/50)
AAWP14250	0.80um MF-Membrane,
AP1512450	142mm (pk/50) Glass Fiber Prefilter,
AP2512450	124mm, "Fine" (pk/50) Glass Fiber Prefilter,
	124mm, "Coarse" (pk/50)

ACCESSORIES

T.C.Gasket, 1 1/2*	10/pk	YY 20 0
Leg Cap, Rubber	3/pk	YY221
O-Ring, Viton®	4/pk	YY22:
O-Ring, LCR-treated Buna-N	5/pk	YY401
PVC Tubing with Clamp, 1/2" X 10	,	XX670

ASSEMBLY OF EQUIPMENT

- 1. Remove handwheel bolts from top of unit and remove top-plates of holder assembly. Install Teflon O-ring in the groove at the base of the stainless steel cylinder. Examine under-drain screen to ensure that the radial spokes are down. Then replace support screen.
- Weigh filters and prefilters to be used or use average weights of 142 Million filters listed below:

Filter	Prefilter
AA - 0.73g	AP25 - 2.64g
DA - 0.79g	AP15 - 1.04g
HA - 0 88a	

Wet filter(s) by floating in pan of deionized water.

- 3. Place required filter(s) on base support screen. If more than one filter is required, place HA filter on screen first, followed by the other filter(s) increasing pore size (i.e. DA, AA).
- 4. Center the stainless steel cylinder (grooved end down) onto the membrane(s) assure that an even seal is obtained.
- 5. Install the AP15 prefilter (if needed) followed by the AP25 prefilter are go sides up) by lowering through the cylinder top onto the membrane(s). The AP prefilter may appear oversized, however, it will abut the sides of the cylinder.
- 6. Place top plate of the filter holder on top cylinder rim. The red silicone O-ring fitted into groove of plate should seal evenly onto rim. Replace handwheel bolts to top of unit and screw down evenly and snugly until wrist tight. Additional tightening must be accomplished by means of plastic write provided.
- 7. Fit the PVC tubing over the Tri-Clover*hose adaptor and secure with hose la Attach other end of PVC tubing to a nitrogen tank regulator outlet and secur with a hose clamp. Put aside until you are ready for step (2) under "Sam le Preparation and Filtration Procedure".

SAMPLE PREPARATION AND FILTRATION PROCEDURE

- 1. Take a representative sample (minimum size 100g) of the waste to be tested
- 2. Using a funnel, pour test material to be filtered into the unit through the Tri-Clamp opening. Attach the TC hose adaptor (fitted with 1 1/2" gasket the TC connection of top plate, using the stainless steel TC clamp provide
- 3. Place appropriate receiving vessel below TC sanitary outlet or attach PVC to TC adaptor and run hose to appropriate container.
- 4. Slowly increase pressure on nitrgen tank. When liquid flow begins from our of assembled holder, immediately turn regulator valve handle a few turns to lower pressure. Bleed air from top-plate vent valve slowly until pressure gauge reads between 10-15 psi. Close vent valve and (if necessary) readjuately regulator valve to set pressure between 10-15 psig. Continue filtration of flow ceases.
- 5. Increase inlet pressure stepwise in 1C psi increments to 75 psi and contin filtration until flow ceases. No leaks should be apparent.
- 6. Shut off pressure from nitrogen tank regulator outlet valve, then open ver valve gradually to release pressure from cylinder.
- 7. Remove inlet TC hose connector and handwheel bolts, then remove top plate extension cylinder of holder.
- 8. Keep the liquid fraction under refrigeration at 1-5°C (34-41°F) for future analysis. The liquid should be clear of particulate material.
- 9. Remove filters, prefilters, and solid cake material. Prepare the solid material for extraction by either grinding it (including filters and prefilters) to pass through a 9.5 mm (3.8") standard serve or by subjectito the structural integrity procedure outlined in the Federal Register.
 - *Federal Register, Dec. 18, 1978. Vol. 43, No. 243.

- 10. Weigh the ground solid material and subtract from this the weight of the filters and prefilters that were used, (Step No. 2 under Assembly of Equipment). Measure a volume of water at 4°C that is equal in weight to material previously collected. The figure obtained is to be applied to the procedure in Step (15) below.
- 11. Place the solid material in an extractor. Information on a suitable extraction can be found in the Federal Register.**
- 12. Add to the extractor a weight of deionized water equal to 16 times the weight of the solid material added to the extractor. This includes any water as a transferring the solid material to the extractor.
- 13. Begin agitation and adjust the pH of the solution to 5.0 \pm 0.2 using 0.5 N acetic acid according to the methodology stated in the Federal Register.*
- 14. At the end of the 24 (± 0.5) hours extraction period, separate the extracte: material into solid and liquid phases by repeating above procedures.
- 15. Adjust the volume of the resulting liquid phase with deionized water so that its volume is twenty times that occupied by a quantity of water at 400 et all weight to the initial quantity of solid material in step (10) above. Combit this solution with the original liquid phase in step (8).
- 16. The analysis conducted to determine conformance with the toxicity criterion must be made with the methods (or equivalent) outlined in the Federal Register.*

^{*}Federal Register, Dec. 18, 1978. Vol. 43, No. 243.

187423

DISCUSSION

Experience has shown that many waste samples (after extraction) can be filtered within 20 minutes using only an AP25 prefilter superimposed on an HA (0.45um) for Some examples include: paper mill effluents, slime muds, secondary sludge, and fash.

Primary effluents, such as fibers, may require a combination of an AP15 prefilte and a DA (0.65um) or AA (0.8um) filter in addition to an HA filter to accomplish adequate filtration. In setting up such combinations the filter with the smalle pore size is placed onto the support screen, followed by other filters from fine coarse with the prefilter always on top.

Difficult-to-filter samples, such as acidified extractions of wood bark, may rec a full complement of filters.

If samples appear cloudy after filtration through an HA filter, they should be filtered through a GS (0.22um pore size) filter.

CAUTION

- Pressure introduced into the system should not exceed 100 psi.
- All flexible tubing used with the system should be able to withstand press exceeding 100 psi.

FOR TECHNICAL ASSISTANCE

Write: Millipore Corporation

Ashby Road

Bedford, Massachusetts, 01730

C/O Technical Service

Call: Millipore Corporation

Technical Service 1-800-225-1380

In Massachusetts: (617) 275-9200

To Place an Order:

Call 800-225-1380 (Toll Free)
And ask for "Order Services".

Millipore is a registered trademark of Millipore Corporation, Bedford, MA. 01730

REV. 4/86

Tri-Clamp and Tri-Clover are trademarks of Ladish Co.

Teflon, and Viton are trademarks of E.I.Dupont de Nemours & Co. Inc.

- d. <u>Collect Sample</u> Groundwater sampling will be conducted by using dedicated bladder pumps or bailers. Both collection procedures are described below.
 - Bladder Pump After purging, the air hose will be fitted to the air line nipple on the bladder pump and the compressor will be turned on. The discharge rate for the bladder pump should be set around 100 mls/minute for successful volatile This rate will be measured by using a 100 ml graduated cylinder and timing the flow. flow rate will be adjusted with the controller until 100 mls/minute is achieved. Volatile samples will be collected first. Vials for volatile analysis will be completely filled leaving no air Semi-volatile samples 'll be filled next followed by the pesticide/PCBs, herbicides, ions, total metals, pH, specific conductance, temperature, and total dissolved solids. Dissolved metal samples will be collected last by placing a 0.45 micron filter in-line. sample will be The collected into a preserved container.
 - d.b. <u>Bailer</u> After purging the well and sufficient recharge has been allowed, samples for volatile organics should be collected using the first bailer volume. Lower the bailer slowly until it contacts the water surface, and allow the bailer to sink to the desired depth and fill, with a minimum of surface disturbance. Slowly withdraw the bailer, taking care to prevent contact of the bailer line with the ground. Slowly discharge the contents into the appropriate sample containers. Repeat the process as necessary to fill each

container to the required volume. Vials for volatile analysis will be completely filled, leaving no air space above the liquid portion (to minimize volatilization). Check that the Teflon on the Teflon-lined silicone septum is toward the sample in the caps and secure the cap tightly. semi-volatile compounds are to be sampled for, collect these samples next. Proceed to the collection of samples for the remaining analyses. Be careful of all pre-preserved bottles. If acids are present, open the bottle upwind and away from the body. All samples for dissolved metals should be collected last and filtered through a 0.45 um filter within 15 minutes after collection. Millipore filtration system will be utilized for this procedure. The procedure for Millipore filtering is as f 'lows:

- 1. Assemble the Millipore filtration unit. Assembly procedures can be found in Appendix C.
- 2. Using a funnel, pour test material to be filtered into the unit through the Tri-Clamp opening. Attach the TC hose adaptor (fitted with 1-1/2" gasket) to the TC connection of top plate, using the stainless steel TC clamp provided.
- 3. Place appropriate receiving vessel below TC sanitary outlet or attach PVC hose to TC adaptor and run hose to appropriate container.
- 4. Slowly increase pressure on nitrogen tank. When liquid flow begins from outlet of assembled holder, immediately turn regulatory valve handle a few turns to lower pressure. Bleed air from top-plate vent valve slowly

until pressure gauge reads between 10-15 psi. Close vent valve and (if necessary) readjust regulator valve to set pressure between 10-15 psi. Continue filtration until flow ceases.

- 5. Increase inlet pressure stepwise in 10 psi increments to 75 psi and continue filtration until flow ceases. No leaks should be apparent.
- 6. Shut off pressure from nitrogen tank regulator outlet valve, then open vent valve gradually to release pressure from cylinder.
- 7. Remove inlet TC hose connector and handwheel bolts, then remove top plate and extension cylinder of holder.
- 8. Acidify e sample with nitric acid to a pH of \leq 2. Test with pH paper.
- 9. Decontaminate the filtration apparatus as outlined in Section 2.3.4.
 - e. <u>Measure Water Level</u> After samples have been collected, the well cap should be replaced. Static water level will be measured again after sufficient recharge has been allowed (24 hours).
 - f. <u>Measure pH. Temperature and Specific Conductance</u> Follow procedure as outlined in Section 2.3.
 - g. <u>Custody</u>, <u>Handling and Shipping</u> Complete the procedures as outlined in Sections 2.2.2 and 2.2.3.

FINAL PAGE

ADMINISTRATIVE RECORD

FINAL PAGE

FINAL PAGE

ADMINISTRATIVE RECORD

FINAL PAGE